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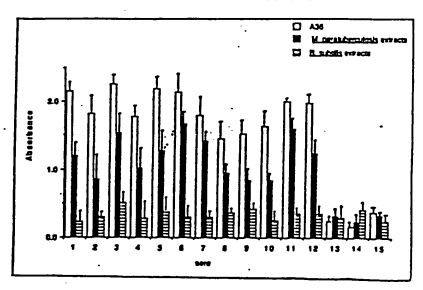
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(54) Title: POLYPEPTIDES FROM MYCOBACTERIUM PARATUBERCULOSIS



(57) Abstract

The invention relates to a polypeptide containing in its polypeptidic chain: the amino acid sequence of 101 amino acids of Figure 8, or a fragment of this sequence, this fragment being such that it is liable to be recognized by antibodies also recognizing the abovesaid sequence of 101 amino acids, but it is not recognized by antibodies respectively raised against M. bovis, M. avium, M. phlei and M. tuberculosis, and possibly against M. leprae, M. intracellulare, M. scrofulaceum, M. fortultum, M. gordonae and M. smegmaths; it is liable to generate antibodies which also recognize the abovesaid sequence of 101 amino acids but which do not recognize M. bovis, M. avium, M. phiel and M. tuberculosis, and possibly M. leprae, M. intracellulare, M. scrofulaceum, M. fortuitum, M. gordonae and M. smegmatis; it reacts with the majority of sera from cattle suffering from Johne's disease; or the polypeptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the above-mentioned properties.

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POLYPEPTIDES FROM MICROBACTERIUM PARATUBERCULOSIS

The invention relates to polypeptides and peptides, particularly recombinant ones, which can be used for the diagnosis of paratuberculosis in cattle and possibly of Crohn's disease in human beings. The invention also relates to a process for preparing the above-said polypeptides and peptides, which are in a state of biological purity such that they can be used as part of the active principle in the preparation of vaccines against paratuberculosis.

It also relates to nucleic acids coding for said polypeptides and peptides.

Purthermore, the invention relates to the <u>in vitro</u> diagnostic methods and kits using the above-said polypeptides and peptides and to the vaccines containing the above-said polypeptides and peptides as active principle against paratuberculosis.

By "recombinant polypeptides or peptides" it is to be understood that it relates to any molecule having a polypeptidic chain liable to be produced by genetic engineering, through transcription and translation, of a corresponding DNA sequence under the control of appropriate regulation elements within an efficient cellular host. Consequently, the expression "recombinant polypeptides" such as is used herein does not exclude the possibility for the polypeptides to comprise other groups, such as glycosylated groups.

The term "recombinant" indeed involves the fact that the polypeptide has been produced by genetic engineering, particularly because it results from the expression in a cellular host of the corresponding nucleic acid sequences which have previously been introduced into an expression vector used in said host.

Nevertheless, it must be understood that the polypeptides or the peptides of the invention can be produced by a different process, for instance by classical chemical synthesis according to methods used in the protein synthesis or by proteolytic cleavage of larger molecules.

The expression "biologically pure" or "biological purity" means on the one hand a grade of purity such that the polypeptides can be used for the production of vaccinating compositions and on the other hand the absence of contaminants, more particularly of natural contaminants.

(Johne's disease) has Paratuberculosis described as one of the most serious diseases affecting the world cattle industry. This mycobacteriosis produced by M. paratuberculosis is characterized by an ileocoecal enteritis leading. successively emaciation, dysentery, cachexy and death (Chiodini R.J. 1984, "Ruminant paratuberculosis (Johne's disease): the current status and future prospects", Cronell Vet. 74:218-262). Histological examination shows oedema, infiltration and thickening of the ileal mucosa, and hypertrophy and necrosis of intestinal lymphnodes. A miliary syndrome with diffused parenchima granuloma in liver, spleen and lungs is not infrequent. The high contagiousness of this disease is due to excretion of large numbers of bacteria from the intestinal tract: contaminated pastures propagate the rapidly producing live-stocks wherein infection, infected animals represent a large part of population. Chronical dysentery is an advanced stage of the disease, for epidemiological data suggest that the subclinical cases, with little sign of intestinal alteration correspond to the majority of infected

animals and frequently to a large proportion of a live-stock population.

paratuberculosis Diagnosis of is especially in the absence of clinical symptoms: it leads to identification of hidden bacterial shedders and avoids propagation of infection. Unfortunately, diagnostic indicators for early stages of the disease are missing. In fact, identification of the etiological agent (a slow grower) is a lengthy process, histological examination material of biopsy difficult and expensive. More interesting appear to be the immunological procedures for analysis of humoral reactions (Brugère-Picoux J., 1987, diagnostic de la paratuberculose chez les ruminants", Rec. Méd. Vét. 163:539-546 ; Colgrave J.S. et al., 1989, "Paratuberculosis in cattle: a comparison of three serologic tests with results of fecal culture", Veterinary Microbiology 19:183-187). complement fixation and hemagglutination apparently lack both sensitivity and specificity, immunoenzymometric methods for evaluation antimycobacterial antibodies seem to be more promising (Abbas B. et al., 1983, "Isolation of specific peptides from <u>Mycobacterium</u> <u>paratuberculosis</u> protoplasm and their use in an enzyme linked immunosorbent assay for the detection of paratuberculosis (Johne's disease) in cattle", Am. J. Vet. Res. 44:2229-2236 ; Colgrave J.S. et al., 1989, "Paratuberculosis in cattle: a comparison of three serologic tests with results of fecal culture" Veterinary Microbiology, 19:183-187; Yokomizo Y. et 1983, "Enzyme-linked immunosorbent assay for detection of bovine immunoglobulin G1 antibody to a protoplasmic antigen of Mycobacterium paratuberculosis" Am. J. Vet. Res. 44:2205-2207; Yokomizo Y. et al., 1985, "A method for avoiding false-positive reactions in an enzyme-linked immunosorbent assay (ELISA) for the

diagnosis of bovine paratuberculosis Japan, J. vet. Sci. 47:111-119).

Moreover, since slaughtering of cattle affected by tuberculosis (caused by M. bovis and/or tuberculosis), but not of those with paratuberculosis, is compulsory in Occidental countries, a distinction at the immunological level between the two mycobacterial diseases is essential. Moreover, M. paratuberculosis is known to be genetically close-related to M. avium (Chiodini R.J. et al., 1989, "The genetic relationship between Mycobacterium paratuberculosis and the M. avium complex" Acta Leprol. 7:249-251; Hurley S.S. et al., "Deoxyribonucleic 1988, acid-relatedness Mycobacterium paratuberculosis to others members of the family Mycobacteriaceae" Int. J. Syst. Bacteriol. 38:143-146), which is a possible host of the intestinal tract of ruminants.

Taking into account the cross reactivity between M. paratuberculosis and many other mycobacteria, it was a priori a difficult approach to find an antigen containing specific epitopes liable to be used as reagents for the diagnosis of paratuberculosis, said reagents having no cross reactivity with other close related mycobacteria.

In addition to the above-mentioned aspects relative to paratuberculosis in cattle, <u>M. paratuberculosis</u> has been found to play an etiologic role in at least some cases of Crohn's disease in human.

The disease originally described by Crohn and coworkers was chronical ileitis producing hyperplastic granulomata of the intestine lymphnodes. The syndrome presently known as Crohn's disease entails inflammatory alterations of different organs of the digestive tract (month, esophagus, stomach, ileum and colon). Segments of the

motive apparatus (joints, muscles and bones) can also be involved. Isolation of mycobacteria from patients affected by the Crohn's disease has been repeatedly related: in several instances isolates were identified as M. paratuberculosis. The induction by these isolates of a syndrome mimicking Crohn's disease in laboratory animals and primates has been successful. In a recent review article (Chiodini R.J., 1989, "Crohn's disease and the mycobacterioses: a review and comparison of two disease entities", Clin. Microbiol. Rev. 2:90-117), Chiodini suggests this syndrome to be the expression of several pathological entities and concludes, that, if Crohn's disease has a mycobacterial etiology, the most likely agent would be M. paratuberculosis.

At this present time, larger epidemiological investigation with an ELISA based on a specific protein of <u>M. paratuberculosis</u> is expected to help to solve the problem of the etiology of this enteritis resembling in many respects the Johne's disease of cattle.

The expression "cattle" means ruminants, such as bovines, sheeps, goats, cervidae, but also include some non ruminant animals which may also be infected by Johne's disease such as monkeys and horses.

An aspect of the invention is to provide recombinant polypeptides which can be used as purified antigens for the detection and control of paratuberculosis.

Another aspect of the invention is to provide nucleic acids coding for the peptidic chains of biologically pure recombinant polypeptides which enable their preparation on a large scale.

Another aspect of the invention is to provide antigens which can be used in serological tests as an <u>in vitro</u> rapid diagnosis of paratuberculosis, as well as in skin tests for <u>in vivo</u> diagnosis of

paratuberculosis and as an immunogenic principle in vaccines.

Another aspect of the invention is to provide a rapid in vitro diagnostic means for paratuberculosis, enabling it to discriminate between cattle suffering from tuberculosis from the ones suffering from paratuberculosis.

Another aspect of the invention is to provide a rapid in vitro diagnostic means for paratuberculosis, enabling it to discriminate between cattle suffering from paratuberculosis from the ones infected or colonized by M. avium, M. bovis or M. tuberculosis or M. phlei.

Another aspect of the invention is to provide <u>in</u> <u>vitro</u> diagnostic means for patients suffering from Crohn's disease.

The invention relates to an antigen complex from M. paratuberculosis, named hereafter "the antigen A36", liable to be obtained as follows:

- sonication of bacterial suspensions of \underline{M} .

 paratuberculosis to obtain a homogenate (also named sonicate),
- centrifugation of the above-mentioned homogenate to obtain a supernatant (which corresponds to the cytoplasm of the bacteria),
- RNAase digestion of the above-mentioned supernatant,
- fractionation of the digested supernatant, for instance by gel exclusion chromatography, for instance on Sepharose 6B columns,
- recovery of the antigen complex (A36) which is the excluded fraction of the fractionation.

It is to be noted that the antigen complex hereabove defined corresponds to the TMA complex (thermostable macromolecular antigens), belonging to a family of complexes present in all mycobacteria and

consisting of or containing lipid, polysaccharide and protein moieties.

The proteic part of the antigen complex of the invention can be fractionated and visualized as follows:

- fractionation of the proteins of the above-mentioned antigen complex by electrophoresis in a gel, for instance 10% polyacrylamide gels to obtain the protein on bands,
- detection of the proteins by staining for instance with Coomassie blue.

The polypeptides of the invention contain in their polypeptidic chain:

- the amino acid sequence of 101 amino acids of Figure 8,
- or a fragment of this sequence, this fragment being such that:
 - . it is liable to be recognized by antibodies also recognizing the abovesaid sequence of 101 amino acids, but it is not recognized by antibodies raised respectively against <u>M. bovis</u>, <u>M. avium</u>, <u>M. phlei</u> and <u>M. tuberculosis</u>,
 - . it is liable to generate antibodies which also recognize the abovesaid sequence of 101 amino acids but which do not recognize M. bovis, M. avium, M. phlei and M. tuberculosis,
 - . it reacts with the majority of sera from cattle suffering from Johne's disease,
- or the polypeptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the above-mentioned properties.

Recognition of one of the above-mentioned fragments by the above-mentioned antibodies - or of the abovesaid sequence of 101 amino acids by the above-

mentioned antibodies - means that the above-mentioned fragment can form a complex with one of the above-said antibodies.

The formation of the complex antigen (i.e. the sequence of 101 amino acids or of the above-said fragment) - antibody and the detection of the existence of a formed complex can be done according to classical techniques such as the ones using a marker labeled by radioactive isotopes or by an enzyme.

Hereafter is also given in a non limitative way, a test for giving evidence of the fact that polypeptides of the invention are recognized selectively by the majority of the sera from cattle suffering from Johne's disease (immunodominant polypeptides), for instance bovines.

This test is an immunoblotting (Western blotting) analysis, in the case where the polypeptides of the invention are obtained by recombinant techniques. This test can also be used for polypeptides of the invention obtained by a different preparation process. After sodium dodecvl sulfate-polyacrylamide gel electrophoresis, polypeptides of the invention are blotted onto nitrocellulose membranes (Hybond (Amersham)) as described by Towbin H. et al., 1979, "Electrophoretic transfer of pioteins polyacrylamide gels to nitrocellulose sheets: procedure and some applications", Proc. Natl. Acad. Sci. USA 76:4350-4354. The expression of polypeptides of the invention fused to β -galactosidase in E. coli Y1089, is visualized by the binding of a polyclonal rabbit anti-A36 antiserum (or polyclonal rabbit anti-homogenate antiserum defined hereafter in the examples, polyclonal rabbit anti-βgal-p362 antiserum, defined hereafter in the examples) (1:1,000) or by using a monoclonal anti- β -galactosidase antibody (Promega). The secondary antibody (anti-rabbit immunoglobulin G and

anti-mouse immunoglobulin G respectively, both alkaline phosphatase conjugated) is diluted as recommended by the supplier (Promega). Colour reaction is developed by adding NBT/BCIP (Nitro Blue Tetrazolium 5-bromo 4-chloro-3-indolyl phosphate [Promega]) using conditions recommended by suppliers.

In order to identify selective recognition of polypeptides of the invention and of fusion proteins of the invention by sera of bovine suffering from Johne's disease, nitrocellulose sheets are incubated overnight with each of these sera (1:50) (after blocking aspecific protein-binding sites).

Reactive areas on the nitrocellulose sheets are revealed by incubation with peroxidase conjugated goat anti-bovine immunoglobulin G antibody (Dakopatts, Copenhagen, Denmark) (1:200) for 4h, and after repeated washings, color reaction is developed by adding α -chloronaphtol (Bio-Rad Laboratories, Richmond, Calif.) in the presence of hydrogen peroxide.

The non-recognition of the antibodies raised against the above-mentioned fragments of the invention by M. bovis, M. avium, M. phlei and M. tuberculosis and by other mycobacteria can be done according to a process detailed in the examples.

As to the non-recognition of the above-mentioned fragments of the invention by antibodies raised respectively against <u>M. bovis</u>, <u>M. avium</u>, <u>M. phlei</u> and <u>M. tuberculosis</u> or other mycobacteria, it can also be done according to a process detailed in the examples.

Advantageous above-defined fragments of the invention are liable not to be recognized by antibodies raised against other mycobacteria such as M. leprae, M. intracellulare, M. scrofulaceum, M. fortuitum, M. gordonae and M. smeqmatis, and are liable to generate antibodies which do not recognize M. leprae, M.

intracellulare, M. scrofulaceum, M. fortuitum, M. gordonae and M. smegmatis.

It goes without saying that the free reactive functions which are present in some of the amino acids, which are part of the constitution of the polypeptides of the invention, particularly the free carboxyl groups which are carried by the groups Glu and Asp or by the C-terminal amino acid on the one hand and/or the free NH₂ groups carried by the N-terminal amino acid or by amino acids inside the peptidic chain, for instance Lys, on the other hand, can be modified in so far as this modification does not alter the above mentioned properties of the polypeptide.

The molecules which are thus modified are naturally part of the invention. The above mentioned carboxyl groups can be acylated or esterified.

modifications Other are also part of invention. Particularly, the amine or carboxyl functions or both of terminal amino acids can be themselves involved in the bond with other amino acids. For instance, the N-terminal amino acid can be linked the C-terminal amino acid of another peptide comprising from 1 to several amino acids.

Furthermore, any peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids of the polypeptides according to the invention are part of the invention in so far as this modification does not alter the above mentioned properties of said polypeptides.

The polypeptides according to the invention can be glycosylated or not, particularly in some of their glycosylation sites of the type Asn-X-Ser or Asn-X-Thr, X representing any amino acid.

An advantageous recombinant polypeptide of the invention is constituted by the sequence represented on Figure 8, extending from the extremity constituted by

amino acid at position (1) to the extremity constituted by amino acid at position (101), or by the following peptides:

Glu-Phe-Pro-Gly-Gly-Gln-His-Ser-Pro-Gln,

(position 1 to 11 on Figure 8)

Gln-Gln-Ser-Tyr-Gly-Gln-Glu-Pro-Ser-Ser-Pro-Ser-Gly-Pro-Thr-Pro-Ala

(position 85 to 101 on Figure 8).

It is to be noted that this polypeptide is derived from the expression product of a DNA derived from the nucleotide sequence coding for a polypeptide of 10 kDa being the carboxy terminal part of a 34 kDa protein of M. paratuberculosis, defined hereafter.

An advantageous recombinant polypeptide of the invention is characterized by the fact that:

- it contains the amino sequence of 101 amino acids of Pigure 8 as its C-terminal part,
- it has a molecular weight of about 34kDa, in SDS-PAGE,
- it is coded by a nucleotide sequence liable to hybridize with the complementary strand of the sequence of Figure 11,
- it reacts with the majority of sera from cattle suffering from Johne's disease,
- it is advantageously liable to elicit a cellular immune response in sensitized subjects.

Subjects can be either test animals such as mice or guinea pigs or cattle or human beings.

"Sensitized" means that these subjects have been in contact previously with <u>M. paratuberculosis</u>, resulting in a priming of the cellular immune system.

Sensitization can be induced by inoculating the subjects with killed or attenuated \underline{M} . paratuberculosis or it can result from a natural infection with \underline{M} . paratuberculosis.

A positive cellular immune response to the polypeptides of the invention can be detected for example in vivo by a delayed - type hypersensitivity reaction upon skintesting with the polypeptides of the invention or in vitro by proliferation of peripheral blood lymphocytes isolated from sensitized subjects, in response to the added polypeptides.

An advantageous recombinant polypeptide of the invention contains or is constituted by the amino acid sequence of Figure 11.

Another advantageous recombinant polypeptide of the invention contains or is constituted by the amino acid sequence extending from amino acid at position (1) to the amino acid at position (199), of Figure 11.

It is to be noted that this polypeptide is a 34 kDa protein which is present in the proteic part of the TMA complex of \underline{M} . paratuberculosis (A36).

Hereafter is given, in a non limitative way, a process for preparing this 34 kDa protein of the invention.

The DNA sequence (306 bp) coding for p362, being the carboxyterminal end of the 34 kDa protein has been determined (see Figure 8). It contains a unique ApaI (GGGCCC) site at position 141.

Using this information, the full gene coding for - the 34 kDa protein can be isolated as follows:

An oligonucleotide coding for a stretch of at least 30 bp, situated within the region EcoRI-ApaI (1-141 bp) of the known sequence, is synthesized.

It is labeled and used as a probe to hybridize to the DNA of M. paratuberculosis (strain ATCC 19698), which has previously been cut by ApaI, separated by agarose gel electrophoresis, denatured and transferred to a nylon membrane.

This hybridization indicates a band on the nylon membrane of around 1500 bp, which contains the coding

part for the rest of the 34 kDa protein. After having located this 1500 bp fragment, flanked by 2 ApaI sites, in the agarose gel, it is isolated from the gel, purified and subcloned in the ApaI site of the sequencing vector pBluescript SK.

After sequencing of this fragment, the coding region, starting with the initiation codon ATG or GTG, is delineated. Using a restriction site near the initiation codon (5' end), naturally present or created by site-directed mutagenesis, and the ApaI site at the 3' end, the DNA fragment coding for the N-terminal part of the protein (about 750 bp) is excised from pBluescript SK', and purified. It is ligated to the ApaI site of the fragment coding for the C-terminal part of p362 (142-306, Figure 8), that for example has been prepared synthetically.

The complete gene coding for the 34 kDa protein (about 910 bp) is subcloned in an expression vector and expressed in <u>E. coli</u>. The recombinant 34 kDa protein is then purified.

The invention also relates to the amino acid sequences constituted by the above mentioned polypeptides and a protein or an heterologous sequence with respect to said polypeptide, said protein or heterologous sequence comprising for instance from about 1 to about 1100 amino acids. These amino acid sequences will be called fusion proteins.

In an advantageous fusion protein of the invention, the heterologous protein is β -galactosidase.

The invention also relates to a nucleic acid characterized by the fact that it comprises or is constituted by:

- a nucleotide chain liable to hybridize with the nucleotide chain coding for the polypeptides according to the invention, or

- a nucleotide chain coding for the polypeptides according to the invention, or
- the complementary sequences of the above nucleotide chains.

The invention also relates to nucleic acids comprising nucleotide sequences which hybridize with the nucleotide sequences coding for any of the above mentioned polypeptides under the following hybridization conditions:

- hybridization and wash medium:
- * a preferred hybridization medium contains about 3 x SSC [SSC = 0.15 M sodium chloride, 0.015 M sodium citrate, pH 7], about 25 mM of phosphate buffer pH 7.1, and 20% deionized formamide, 0.02% Ficoll, 0.02% BSA, 0.02% polyvinylpyrrolidone and about 0.1 mg/ml sheared denatured salmon sperm DNA,
- * a preferred wash medium contains about 3 x SSC, about 25 mM phosphate buffer, pH 7.1 and 20% deionized formamide;
- hybridization temperature (HT) and wash temperature (WT) for the nucleic acids of the invention defined by x-y: i.e. by the sequence extending from the extremity constituted by the nucleotide at position (x) to the extremity constituted by the nucleotide at position (y) represented on Figures 7A, 7B or 7C:
- 1 306 (for Figures 7B and 7C) or

HT = WT = 65°C

- 1 307 (for Figure 7A)
- 1 507 (for Figures 7B and 7c)

HT = WT = 65°C

1 - 508 (for Figure 7A)

The above mentioned temperatures are to be considered as approximately ± 5°C.

It is to be noted that in the above defined nucleic acids, as well as in the hereafter defined

nucleic acids, the nucleotide sequences which are brought into play are such that T can be replaced by U.

A group of preferred nucleic acids of the invention comprises one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (307) represented in Figure 7A,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (508) represented in Figure 7A, wherein
- X and E represent phosphodiester bonds,
- Y and F represent respectively G and C,
- Z and H represent respectively C and G, or
- X and E represent respectively G and C,
- Y and P represent respectively C and G,
- Z and H represent phosphodiester bonds.

A group of preferred nucleic acids of the invention comprises one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (306) represented in Figure 7B,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (507) represented in Figure 7B.

The nucleotide sequence represented in Figure 7B corresponds to the one represented in Figure 7A, wherein

- X and E represent phosphodiester bonds,
- Y and F represent respectively G and C,

- Z and H represent respectively C and G.

The invention also relates to a nucleic acid characterized by the fact that it comprises or is constituted by a nucleotide chain,

- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (306) on Figure 7C, or
- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (507) on Figure 7C.

The nucleotide sequence represented on Figure 7C corresponds to the one represented on Figure 7A, wherein

- X and E represent respectively G and C,
- Y and F represent respectively C and G,
- Z and H represent phosphodiester bonds.

The invention also relates to a nucleic acid which comprises or is constituted by:

- a nucleotide sequence liable to hybridize with the complementary strand of the nucleotide sequence of Figure 11, or with the complementary strand of the nucleotide sequence extending from nucleotide at position (742) to nucleotide at position (1338) of Figure 11,
- the nucleotide sequence of Figure 11 or the nucleotide sequence extending from nucleotide at position (742) to nucleotide at position (1338) of Figure 11,
- the complementary sequences of the nucleotide sequences above-defined.

From the nucleic acids of the invention, probes (i.e. cloned or synthetic oligonucleotides) can be inferred.

These probes can be from 15 to the maximum number of nucleotides of the selected nucleic acids. The oligonucleotides can also be used either as

amplification primers in the PCR technique (PCR, Mullis and Falcona, Methods in Enzymology, vol. 155, p. 335, 1987) to generate specific enzymatically amplified fragments and/or as probes to detect fragments amplified between bracketing oligonucleotide primers.

The specificity of a PCR-assisted hybridization assay can be controlled at different levels.

The amplification process or the detection process or both can be specific. The latter case giving the higher specificity is preferred.

The invention also relates to any recombinant nucleic acid containing at least one of the nucleic acids of the invention combined to or inserted in a heterologous nucleic acid.

The invention relates more particularly to recombinant nucleic acid such as defined, in which the nucleotide sequence of the invention is preceded by a promoter (particularly an inducible promoter) under the control of which the transcription of said sequence is liable to be processed and possibly followed by a sequence coding for transcription termination signals.

The invention also relates to the recombinant nucleic acids in which the nucleic acid sequences coding for the polypeptide of the invention and possibly the signal peptide, are recombined with control elements which are heterologous with respect to the ones to which they are normally associated with in the mycobacterial genome and, more particularly, the regulation elements adapted to control their expression in the cellular host which has been chosen for their production.

The invention also relates to recombinant vectors, particularly for cloning and/or expression, comprising a vector sequence, notably of the type plasmid, cosmid or phage or virus DNA, and a recombinant nucleic acid

of the invention, inserted in one of the non essential sites for its replication.

According to an advantageous embodiment of the invention, the recombinant vector contains necessary elements to promote the expression in a cellular host of polypeptides coded by nucleic acids according to the invention inserted in said vector and notably a promoter recognized by the RNA polymerase of the cellular host, particularly an inducible promoter and possibly a sequence coding for transcription termination signals and possibly a signal sequence and/or an anchoring sequence.

According to another additional embodiment of the invention, the recombinant vector contains the elements enabling the expression by \underline{E} . coli of a fusion protein consisting of the polypeptide of β -galactosidase or part thereof linked to a polypeptide coded by a nucleic acid according to the invention.

The invention also relates to a cellular host, chosen from among bacteria such as <u>E. coli</u> or chosen from among eukaryotic organism, such as CHO cells or insect cells, which is transformed by a recombinant vector according to the invention, and containing the regulation elements enabling the expression of the nucleotide sequence coding for the polypeptide according to the invention in this host.

The invention relates to an expression product of a nucleic acid expressed by a transformed cellular host according to the invention.

The invention also relates to a process for preparing a recombinant polypeptide according to the invention comprising the following steps:

- the culture in an appropriate medium of a cellular host which has previously been transformed by an appropriate vector containing a nucleic acid according to the invention,

- the recovery of the polypeptide produced by the above said transformed cellular host from the above said culture medium, or from the cellular host, and
 - possibly the purification of the polypeptide produced, eventually by means of immobilized metal ion affinity chromatography (DMAC).

The polypeptides of the invention can be prepared according to the classical techniques in the field of peptide synthesis.

The synthesis can be carried out in homogeneous solution or in solid phase.

For instance, the synthesis technique in homogeneous solution which can be used is the one described by Houbenweyl in the book titled "Methode der organischen chemie" (Method of organic chemistry) edited by E. Wunsh, vol. 15-I et II. THIEME, Stuttgart 1974.

The polypeptides of the invention can also be prepared in solid phase according to the method described by Atherton & Shepard in their book titled "Solid phase peptide synthesis" (Ed. IRL Press, Oxford, NY, Tokyo, 1989).

The invention also relates to a process for preparing the nucleic acids according to the invention.

A suitable method for chemically preparing the single-stranded nucleic acids (containing at most 100 nucleotides of the invention) comprises the following steps:

- DNA synthesis using the automatic β -cyanoethyl phosphoramidite method described in Bioorganic Chemistry 4; 274-325, 1986.

In the case of single-stranded DNA, the material which is obtained at the end of the DNA synthesis can be used as such.

A suitable method for chemically preparing the double-stranded nucleic acids (containing at most

100 bp of the invention) comprises the following steps:

- DNA synthesis of one sense oligonucleotide using the automatic β -cyanoethyl phosphoramidite method described in Bioorganic Chemistry 4; 274-325, 1986, and DNA synthesis of one anti-sense oligonucleotide using said above-mentioned automatic β -cyanoethyl phosphoramidite method,
- combining the sense and anti-sense oligonucleotides by hybridization in order to form a DNA duplex,
- cloning the DNA duplex obtained into a suitable plasmid vector and recovery of the DNA according to classical methods, such as restriction enzyme digestion and agarose gel electrophoresis.

A method for the chemical preparation of nucleic acids of length greater than 100 nucleotides - or bp, in the case of double-stranded nucleic acids - comprises the following steps:

- assembling of chemically synthesized oligonucleotides, provided at their ends with different restriction sites, the sequences of which are compatible with the succession of amino acids in the natural peptide, according to the principle described in Proc. Nat. Acad. Sci. USA 80; 7461-7465, 1983,
- cloning the DNA thereby obtained into a suitable plasmid vector and recovery of the desired nucleic acid according to classical methods, such as restriction enzyme digestion and agarose gel electrophoresis.

The invention also relates to antibodies themselves formed against the polypeptides according to the invention, and characterized by the fact that they recognize neither <u>M. bovis</u>, nor <u>M. avium</u>, nor <u>M. phlei</u>, nor <u>M. tuberculosis</u>.

It goes without saying that this production is not limited to polyclonal antibodies.

It also relates to any monoclonal antibody produced by any hybridoma liable to be formed according to classical methods from splenic cells of an animal, particularly of a mouse or rat, immunized against the purified polypeptide of the invention on the one hand, and of cells of a myeloma cell line on the other hand, and to be selected by its ability to produce the monoclonal antibodies recognizing the polypeptide which has been initially used for the immunization of the animals.

The invention also relates to any antibody of the invention labeled by an appropriate label of the enzymatic, fluorescent or radioactive type.

The polypeptide which is advantageously used to produce antibodies, particularly monoclonal antibodies, is the one or part of the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (101) represented on Figure 8.

Variations of this peptide are also possible depending on its intended use. For example, if the peptide is to be used to raise antisera, the peptide may be synthesized with an extra cysteine residue added. This extra cysteine residue is preferably added to the amino terminus and facilitates the coupling of the peptide to a carrier protein which is necessary to render the small peptide immunogenic. If the peptide is to be labeled for use in radioimmune assays, it may be advantageous to synthesize the protein with a tyrosine attached to either the amino or carboxyl terminus to facilitate iodination. This peptide possesses therefore the primary sequence of the peptide above-mentioned but with additional amino acids which do not appear in the primary sequence of the protein and whose sole function is to confer the desired chemical properties to the peptide.

The invention also relates to a process for detecting <u>in vitro</u> antibodies related to paratuberculosis in a biological sample of an animal liable to contain them, this process comprising

- contacting the biological sample with a polypeptide or a peptide according to the invention, or the expression product of the invention, under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and
- the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

Preferably, the biological medium is constituted by an animal serum, and particularly by bovine serum.

The detection can be carried out according to any classical process.

By way of example a preferred method brings into play an immunoenzymatic process according to ELISA technique or immunofluorescent or radioimmunological (RIA) or the equivalent ones.

Thus the invention also relates to any polypeptide according to the invention labeled by an appropriate label of the enzymatic, fluorescent, radioactive... type.

Such a method for detecting <u>in vitro</u> antibodies related to paratuberculosis comprises for instance the following steps:

- deposit of determined amounts of a polypeptidic composition according to the invention in the wells of a titration microplate,
- introduction into said wells of increasing dilutions of the serum to be diagnosed,
- incubation of the microplate,
- repeated rinsing of the microplate,
- introduction into the wells of the microplate of labeled antibodies against the blood immunoglobulins,

- the labeling of these antibodies being based on the activity of an entyme which is selected from among the ones which are able to hydrolyze a substrate by modifying the absorption of the radiation of this latter at least at a given wave length,
- detection by comparing with a control standard of the -amount of hydrolyzed substrate.

The invention also relates to a process for detecting and identifying in vitro antigens of \underline{M} . paratuberculosis in an animal biological sample liable to contain them, this process comprising:

- contacting the biological sample with an appropriate antibody of the invention under conditions enabling an in vitro immunological reaction between said antibody and the antigens of <u>M. paratuberculosis</u> which are possibly present in the biological sample and
- the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

Preferably, the biological medium is constituted by serum or faeces, milk or urine, particularly of bovine origin.

Appropriate antibodies are advantageously monoclonal antibodies directed against the abovementioned peptide.

The invention also relates to an additional method for the <u>in vitro</u> diagnosis of paratuberculosis in an animal liable to be infected by <u>Mycobacterium paratuberculosis</u> comprising:

- contacting a biological sample taken from an animal with a polypeptide or a peptide of the invention, or the expression product of the invention, under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide or peptide and the antibodies which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which has possibly been formed.

To carry out the <u>in vitro</u> diagnostic method for paratuberculosis in an animal liable to be infected by <u>Mycobacterium paratuberculosis</u>, the following necessary or kit can be used, said necessary or kit comprising:

- a polypeptide or a peptide according to the invention, or the expression product of the invention,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned polypeptide or peptide is not labeled.

The invention also relates to an additional method for the <u>in vitro</u> diagnosis of paratuberculosis in an animal liable to be infected by <u>M. paratuberculosis</u>, comprising the following steps:

- contacting a biological sample of said animal with an appropriate antibody of the invention under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. paratuberculosis</u> which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

To carry out the <u>in vitro</u> diagnostic method for paratuberculosis in an animal liable to be infected by <u>Mycobacterium paratuberculosis</u>, the following necessary or kit can be used, said necessary or kit comprising:

- an antibody of the invention,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction, said reagent possibly having a label or being liable to be recognized by a labeled reagent, more

particularly in the case where the above-mentioned antibody is not labeled.

An advantageous kit for the <u>in vitro</u> diagnosis of paratuberculosis comprises:

- at least a suitable solid phase system, e.g. a microtiter-plate for deposition thereon of the biological sample to be diagnosed in vitro,
- a preparation containing one of the monoclonal antibodies of the invention,
- a specific detection system for said monoclonal antibody,
- appropriate buffer solutions for carrying out the immunological reaction between a test sample and said monoclonal antibody on the one hand, and the bonded monoclonal antibodies and the detection system on the other hand.

The invention also relates to a kit, as described above, also containing a preparation of one of the polypeptides or peptides of the invention, said antigen of the invention being either a standard (for quantitative determination of the antigen of M. paratuberculosis which is sought) or a competitor, with respect to the antigen which is sought, for the kit to be used in a competition dosage process.

The invention also relates to a method for the <u>in</u> <u>vitro</u> diagnosis of Crohn's disease in a patient liable to be infected by <u>Mycobacterium</u> <u>paratuberculosis</u> comprising the following steps:

- contacting the biological sample with an appropriate antibody according to the invention, under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. paratuberculosis</u> which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which may be formed:

The invention also relates to a method for the <u>in</u> <u>vitro</u> diagnosis of Crohn's disease in a patient liable to be infected by <u>M. paratuberculosis</u>, comprising the following steps:

- contacting a biological sample taken from a patient with a polypeptide or peptide according to the invention, or the expression product of the invention, under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which has been possibly formed.

The invention also relates to a necessary or kit for an <u>in vitro</u> diagnosis method of Crohn's disease in a patient liable to be infected by <u>Mycobacterium paratuberculosis</u>, said necessary or kit comprising:

- an antibody according to the invention,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction said reagents possibly having a label or being liable to be recognized by a labeled reagent, more particularly in the case where the above-mentioned antibody is not labeled.

The invention also relates to a necessary or kit for an <u>in vitro</u> diagnosis method of Crohn's disease in a patient liable to be infected by <u>Mycobacterium paratuberculosis</u> said necessary or kit comprising:

- a polypeptide or a peptide according to the invention, or the expression product of the invention,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or

being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned polypeptide is not labeled.

The invention also relates to an immunogenic composition comprising a polypeptide or a peptide according to the invention, or the expression product of the invention, in association with a pharmaceutically acceptable vehicle.

invention also relates to composition comprising among other immunogenic principles anyone of the polypeptides or peptides of the invention or the expression product of the invention, possibly coupled to a natural protein or to a synthetic polypeptide having a sufficient molecular weight so that the conjugate is able to induce in vivo the production of antibodies neutralizing Mycobacterium paratuberculosis, induce in _ vivo a protective or immune response cellular by activating M. paratuberculosis antigen-responsive T cells.

The invention also relates to a necessary or kit for the diagnosis of prior exposure of an animal to <u>M. paratuberculosis</u>, said necessary or kit containing a preparation of at least one of the polypeptides or peptides of the invention, or the expression product of the invention, with said preparation being able to induce in vivo after being intradermally injected to an animal a delayed type hypersensitivity reaction, at the site of injection, in case the animal has had prior exposure to <u>M. paratuberculosis</u>.

Other characteristics and advantages of the invention will appear in the following examples and the figures illustrating the invention.

LEGENDS TO FIGURES

 Figure 1(1) represents the two-dimensional cross immunoelectrophoresis (CIE) of total cytoplasm (the supernatant fraction obtained after centrifugation of the sonicate) from <u>M. paratuberculosis</u> and Figure 1(2) represents the two-dimensional cross immunoelectrophoresis of the exclusion fraction obtained by gel exclusion chromatography of the same cytoplasm.

In the second dimension (upward in the figure), migration was made in a gel containing rabbit antiserum against the mycobacterial directed Preparations in 1 and 2 contained 10 µg of proteins. figure identifies the TMA complex of M. the exclusion paratuberculosis (A36) present in fraction.

- Figure 2 represents the serological analysis of infected animals with polypeptide p362. Multiwell plates were coated with 4 μg of proteins/well of E. coli-a362 total cytoplasm (white) or E. coli-control total cytoplasm (black). Samples of diluted (1/400) bovine sera previously exhausted by incubation with E. coli-control homogenate (said homogenate and total cytoplasm being obtained in the same way as M. paratuberculosis homogenate and total cytoplasm as described above) were added, followed by washing, incubation with labeled anti-bovine Ig, peroxidase reagents and spectrophotometric reading at 450 nm.

The following sera were used: asymptomatic non-excretory (sample 1), asymptomatic excretory (samples 2 to 13), symptomatic excretory (samples 14 to 24) and healthy bovine (samples 26 to 32).

- Figure 3 represents the serological analysis of infected animals with a A36-based immunoassay.

Multiwell plates were coated with comparable amounts (0.5 μ g total proteins/well) of: M. paratuberculosis total cytoplasm (black), A36 (white) and B. subtilis total cytoplasm (control: hatched). Samples of diluted (1/400) bovine sera previously

exhausted by incubation with <u>B. subtilis</u> homogenate (said homogenate and total cytoplasm being obtained in the same way as <u>M. paratuberculosis</u> homogenate and total cytoplasm as above-described) were added, followed by washing, incubation with labeled antibovine Ig, peroxidase reagents and spectrophotometric reading at 450 nm. The following bovine sera were used:

a) symptomatic-excretory forms of paratuberculosis (samples 1 to 7); b) asymptomatic-excretory forms (samples 8 to 12); and c) healthy cattle (samples 13 to 15). Mean values of absorbance and standard deviations are the results of 4 repeats.

Pigure 4 represents the recognition of different A36 proteins by the sera of infected bovines. A36 proteins from M. paratuberculosis were fractionated by gel electrophoresis and transferred to nitrocellulose. Membranes were incubated with sera from uninfected (lane 8) or infected animals (lanes 4 to 7), either pre-absorbed (lane 7) or not (lanes 4, 5, 6) with a mixture of homogenates of M. avium, M. bovis and M. phlei. Membrane-bound primary Iq were revealed by labeled secondary Ig. Sera of infected animals were as follows: asymptomatic-non excretory (lane 4), asymptomatic-excretory (lane 5), and symptomaticexcretory (lane 6, 7) cases of paratuberculosis. Reference molecular weight standards (lane 1) and A36 proteins (lane 2) were stained by India ink. Reference: proteins immunoblotted with anti-A36 rabbit antiserum (lane 3).

Figure 5 represents the analysis of the size of the polypeptide (p362) fused to β -galactosidase expressed by recombinant clone a362 (hereafter defined). This fusion protein is named β gal-p362.

Lysate proteins of \underline{E} . \underline{coli} Y1089 lysogenized either by standard $\lambda gtll$ (tracks C and E) or by the same phage carrying the insert coding for p362 (clone

a362) (tracks D and F) were fractionated by 7.5% polyacrylamide gel electrophoresis. Tracks C and D and molecular weight standards (tracks A and B) were stained with Coomassie brilliant blue, whereas tracks E and F were treated with rabbit anti-A36 antiserum and stained with peroxydase-labeled anti-rabbit antiserum.

Figure 6 represents the evidence of the belonging of the recombinant polypeptide p362 to the 34 kD protein of the A36 complex.

The TMA complex from M. paratuberculosis was dissociated and its protein components were fractionated by 10% polyacrylamide gel electrophoresis and transblotted to a nitrocellulose membrane. Fractionated proteins were either stained with India ink (track b) or incubated with rabbit anti- β gal-p362 antiserum (track c). Track a: molecular weight standards.

Figure 7A represents the nucleic acid sequence encompassing the nucleic acid sequence of Figure 7B and the one of Figure 7C.

Figure 7B represents a sequence homologous to the one represented on Figure 7C.

Figure 7C represents the base sequence of the \underline{M} . paratuberculosis genomic fragment present in clone a362 and coding for p362.

It should be noted that the two EcoRI sites [GAATTC] present at both ends of the sequence are a result of the cloning strategy and are not naturally present in the genomic sequence.

Figure 8 represents the amino acid sequence and corresponding nucleotide sequence of the recombinant polypeptide p362.

It should be noted that the first two amino acids, corresponding to the EcoRI sites in the DNA sequence, are not naturally present in the native protein, but are a result of cloning.

Figure 9a corresponds to the restriction and genetic map of the pmTNF-MPH plasmid used in Example II for the expression of p362 of the invention in <u>E. coli</u>.

Figure 9b corresponds to the pmTNF-MPH nucleic acid sequence.

On this figure, the origin of nucleotide stretches used to construct plasmid pmTNF-MPH is specified hereafter.

Position

- 1-208: lambda PL containing EcoRI blunt-MboII blunt fragment of pPL(\(\lambda\)) (Pharmacia)
- 209-436: synthetic DNA fragment
- 230-232: initiation codon (ATG) of mTNF fusion protein
- 236-307: sequence encoding AA 2 to 25 of mature mouse TNF
- 308-384: multiple cloning site containing His encoding sequence at position 315-332
- 385-436: HindIII fragment containing <u>E. coli</u> trp terminator
- 437-943: rrnBT₁T₂ containing HindIII-SspI fragment from pKK223 (Pharmacia)
- 944-3474: DraI-EcoRI blunt fragment of pAT₁₅₃
 (Bioexcellence) containing the tetracycline resistance gene and the origin of replication.

Figure 10 represents the complete amino acid sequence of the recombinant polypeptide mTNF-H6-p362. The amino acids 1-26 represent the mTNF part, the amino acids from 27-46 correspond to the polylinker part (H6) and the remaining amino acids (47-147) represent the <u>M. paratuberculosis</u> 10 kDa polypeptide (p362).

Figure 11 represents the DNA sequence containing the nucleic acid coding for the protein of 34 kDa hereabove defined and the corresponding amino acid sequence. Nucleotides are numbered in the right-hand

side margin and amino acids are numbered below the protein sequence.

It is to be noted that the arrow before amino acid 200 corresponds to the third amino acid of Figure 8, since the first two amino acids of Figure 8 are artificial, corresponding to the <u>EcoRI</u> site from cloning.

Table 5 hereafter corresponds to the complete restriction site analysis of pmTNF-MPH.

267 :

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2333

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Table 5

* RESTRICTION-SITE ANALYSIS *

Done on DNA sequence PHTNPHPH.

Total number of bases is: 3474. Analysis done on the complete sequence.

List of cuts by enzyme.

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EXAMPLE I: Purification of the TMA complex of M. paratuberculosis (A36), characterization of the proteic part of A36, identification of the 34 kDa protein and development of A36 based immunoassay:

MATERIALS AND METHODS

Bacteria:

mycobacteria were used: following The paratuberculosis strain 2E and 316F (from Dr. Saxegaard, National Veterinary Institute, Oslo, Norway; Saregaard F. et al., 1985, "Control of paratuberculosis vaccination" goats by (Johne's disease) in 116:439-441); M. avium serotype 4 (from Dr. Portaels, Institute of Tropical Medicine, Antwerpen, Belgium) (Shaefer W.B., 1965, "Serologic identification and classification of the atypical mycobacteria by their agglutination", Am. Rev. Resp. Dis. suppl. 92:85-93); M. bovis strain BCG GL2 (from Dr. Weckx, Pasteur Institute, Brussels, Belgium) and M. phlei strain AM76 (from Dr. M. Desmecht, National Institute for Veterinary Research, Brussels, Belgium). The 168 strain of B. subtilis was used as control ATCC n° 33234.

Preparation of bacterial cytoplasms:

Bacterial suspensions in buffered saline (100 mg wet weight cells/ml 0.15 M NaCl 0.02 M K2HPO4 pH 7.5 containing 10 mM phenylmethylsulfonyl fluoride) were disrupted by sonication (15 min treatment with a 500-W ultrasonic processor, Vibra cell from Sonics and Materials Inc, Danbury, Co USA (3 min sonication for B. subtilis). Homogenates were centrifuged (5000 x g, 10 min, 4°C), and supernatants (i.e. mycobacterial cytoplasms) were stored at -20°C and used as sources of antigens.

Purification of TMA complexes:

The supernatant (about 4.5 mg proteins/ml) was submitted to RNAase digestion (10 μ g enzyme/100 μ g wet

weight bacteria, 30 min, 37°C) and fractionated by gel exclusion chromatography on Sepharose (Pharmacia, Uppsala, Sweden) equilibrated with buffered saline, as previously detailed (Cocito C. et al., 1986, *Preparation and properties of antigen Mycobacterium bovis BCG" Clin. Exp. 66:262-272). TMA complexes (thermostable macromolecular antigen complexes) were found within the excluded fractions (which contained on the average 0.5 mg soluble proteins/ml). Solutions of TMA (with 1 mM phenylmethylsulfonyl fluoride as conservative) were stored at -20°C.

Purity of TMA complexes was checked by crossed immunoelectrophoresis, according to the reference systems (Closs O. et al., 1980, "The antigens of Mycobacterium bovis, strain BCG, studied by crossed immunoelectrophoresis: a reference system" Scand. J. Immunol. 12:249-263; Gunnarsson E. et al., 1979, "Analysis of antigens in Mycobacterium paratuberculosis" Acta Vet. Scand. 20:200-215).

For this purpose agarose gels (1% type 2 agarose from Sigma, St Louis, Mo) on glass plates (5 by 7 cm) were used, the top gel containing 200 µl of rabbit anti-mycobacterial homogenate. Mycobacterial antigen (10 µl of samples containing 0.5 mg TMA/ml) was applied to a corner well and electrophoretic runs were made as described (1 h, 8 V/cm, 15°C in 1st dimension; 3 V/cm, 18 h, 15°C in 2nd dimension). Slants were washed, dried, stained with Coomassie blue and photographed.

Animal sera:

For production of polyclonal antisera, mycobacterial homogenate or TMA preparations (10 μ g soluble proteins/0.5 ml buffered saline emulsified with equal volume of incomplete Freund adjuvant) were repeatedly injected (6 inoculations at 1-week intervals) into rabbits by subcutaneous way.

The antibody titer of the sera was evaluated by an immunoenzymometric procedure (see below).

Here is thus obtained a polyclonal anti-TMA complex antiserum, more particularly anti-A36 antiserum, and a polyclonal anti-homogenate antiserum referred to in the Western blotting test.

Four kinds of sera from bovines either healthy or at different stages of the Johne's disease were used: a) healthy controls with no sign of mycobacterial infection and with negative tests of coproculture and complement fixation; b) asymptomatic non-excretory stage I of the disease (a case which appeared negative at the moment of sampling but became positive later); c) asymptomatic excretory stage II of the disease (positive coproculture with no clinical signs disease); and d) symptomatic excretory stage III of the disease (with positive complement fixation test). These sera were provided by the National Institute Veterinary Research (Dr. M. Desmecht, Brussels, Belgium) and the Center of Veterinary Medicine (Dr. B. Limbourg, Erpent, Belgium).

Electrophoretic fractionation and Western blotting of TMA proteins:

protein moiety of TMA complexes fractionated by electrophoresis on 10% polyacrylamide gels, in the presence of Na dodecyl sulfate (SDS-PAGE procedure) (Laemmli U.K., 1970, "Cleavage of structural during the assembly of the bacteriophage T4" Nature 227:680-695). Protein samples (25 μ g soluble polypeptides in 50 μ l 0.125 mM Tris-HCl pH 6.8 containing 5% w/v SDS, 20% v/v glycerol, 10% V:V β -mercaptoethanol and 0.05% bromophenol blue) were boiled for 5 min and then applied to vertical gel slabs. Molecular weight protein markers (Sigma Chem. Co., St Louis, Mo) were: bovine serum albumin (66 kDa), ovalbumin (45 kDa), glyceraldehyde-3-phosphate

dehydrogenase (36 kDa), carbonic anhydrase (29 kDa), trypsinogen (24 kDa), trypsin inhibitor (20.1 kDa) and α -lactalbumin (14.2 kDa). Electrophoretic runs (4 h, 50 V, 20°C) were made in a vertical unit (LKB, Bromma, Sweden). Protein bands were visualized by staining with Coomassie brilliant blue. Controls of total cytoplasmic proteins were run in parallel with TMA samples.

Electrophoresed proteins were transferred from polyacrylamide gels to nitrocellulose membranes (BA 85, Macherey-Nagel, Germany) by the use of a transblot-unit (217 multiphor 2, LKB, Bramma, Sweden).

Transfer buffer contained 20% methanol, 0.039 M glycine and 0.048 M Tris base pH 8.8, and runs were made at 10 V for 2 h. Transblotted proteins were identified by reaction with a primary antibody (either polyclonal rabbit antiserum [1/1500] or bovine serum [1/100]) and then with a labeled secondary antibody.

Transblotted nitrocellulose sheets were first incubated for 30 min with TBS buffer (0.5 M NaCl, 0.023 M Tris-HCl pH 7.5) containing 3% w/v gelatin and then for 3 h with the primary antibodies diluted with TBST buffer (TBS containing 0.05% v/v Tween 20) and 1% w/v gelatin. After repeated washings with TBST, sheets were incubated for 2 h with secondary IgG (1/400 diluted preparations of peroxydase-labeled anti-rabbit, anti-mouse or anti-cow IgG, Dako, Copenhagen, Denmark), followed by washings with TBST and TBS buffers. A color reaction was developed by addition of a-chloronaphtol (Bio-Rad Laboratories, Richmond, Cal) in the presence of hydrogen peroxide. The color reaction was stopped by washing sheets with distilled water. A similar protocol antigens directly spotted used for was nitrocellulose membranes (dot-blot analysis). Reference samples of transblotted total proteins and molecular weight markers were visualized by India ink staining (10% solution of fount India, Pelikan, Germany, in 0.2

M NaCl, 0.05 M Tris-HCl pH 7.4 containing 0.3% v/v Tween 20) for 30 min (Hancok K. et al., 1983, "India ink staining of proteins on nitrocellulose paper" Anal. Biochem. 133:157-162).

Immunoassay for determination of antimycobacterial Ig:

Multiwell microtiter plates (Microwell Module, Nunc, Denmark) were coated either with purified A36 or cytoplasm paratuberculosis total with M. supernatant) (0.5 μ g soluble proteins/50 μ l 0.05 M Na carbonate buffer pH 9.6/well). Air dry wells were saturated with bovine serum albumin (0.1% w/v BSA in 0.15 M NaCl, 1 h, 37°C). Increasing dilutions of serum to be tested in 0.15 M NaCl 0.02 M Na phosphate buffer pH 7.2 0.005% Tween 80 (PBST buffer) were added (50 μ l/well, 1 h, 37°C), optimal dilutions being identified by checker board titration. Horse-radish peroxydaselabeled swine anti-rabbit, or rabbit anti-cow antiserum (Dako, Copenhagen, Denmark) were added (50 μ l of 1/400 IgG dilution in PBST/well, 1 h, 37°C). Excess reagent was removed by 5 buffer washings. After incubation with the peroxidase reagent (50 μ l per well of a 17 mM Na citrate buffer pH 6.3 containing 0.2% 0-phenylene diamine and 0.015% H_2O_2 , 30 min, 37°C in the dark), the reaction was stopped (50 μ l 2 M H_2SO_4) and samples were spectrometrically measured (Plate reader SLT 210 from Kontron Analytical, U.K.). Results were recorded as ELISA absorbance values (A450mm).

In some experiments, cross-reactive Ig were removed by incubation (18 h, 4°C) with either purified TMA preparations (0.2 mg protein/ml of serum) or bacterial homogenates or intact mycobacteria (equivalents of 2 mg dry weight bacteria/ml of serum). Absorbed preparations were checked by dot-blot trials before application in immunoblot or immunoassay.

Immune electron microscopy:

Suspensions of mycobacteria in water (5 x 107 cells/5 µl) were placed on carbon-formvar 200-mesh copper grids and air dried. Grids were serially incubated with: a) bovine serum albumin (3% solution in buffered saline, 30 min, 37°C); b) anti-TMA complex rabbit antiserum (a 10°3 dilution of Ig in buffered saline with 0.05% Tween 20, 2 h, 37°C); c) sheep antirabbit biotinylated Ig (1/200 dilution of Ig from Amersham, U.K., in buffered saline-Tween, 1 h, 20°C); d) gold-labeled streptavidin (a 1/20 dilution of a preparation from Amersham, U.K.) (Cloeckaert A. et al., 1990, "Identification of seven surface-exposed Brucella membrane proteins by use antibodies: immunogold labeling for electron microscopy and enzyme-linked immunosorbent assay" Infect. Immun. 58:000-000). Grids were analyzed in a transmission electron microscope (Philips CM 10).

RESULTS

Purification of THA complexes and preparation of anti-THA antisera:

The TMA complex of M. paratuberculosis (A36) has been prepared from the total homogenate. Cytoplasm fractionation by gel exclusion chromatography yielded said TMA complex within the exclusion fraction. The immunoelectrophoretic patterns of total cytoplasmic antigens (supernatant) (Figure 1(1)) and of the exclusion fraction (Figure 1(2)) are compared. From these tracings, which were obtained with polyclonal antisera elicited by inoculation of rabbits with whole mycobacterial homogenate, the purity of the A36 preparation can be inferred.

A similar protocol was used for preparation of other antigens of the TMA group from M. avium, M. bovis and M. phlei, which were used for comparative analysis.

The polyclonal antisera corresponding to the TMA complexes have also been prepared. The purity of these Ig preparations was checked by crossed immunoelectrophoresis: using total cell homogenates as antigens in every case, a single immunoprecipitogen line corresponding to the TMA complex was obtained (patterns not shown, mimicking that of Figure 1(2)). It is to be noted that subcutaneous injection of TMA complex preparations invariably induced the synthesis of high titer antisera (ELISA absorbance higher than 2.5 for dilutions at 10⁻⁵), a result which stressed the high immunogenicity of these antigen complexes.

Development of A36-based serological assay for paratuberculosis:

prompted A36 has availability of development of an enzymometric ELISA-type immunoassay for paratuberculosis. Accordingly, multiwell plates were coated with A36 and incubated with sera infected animals. Peroxidase-labeled rabbit anti-bovine IgG were added as second antibody, and the color developed after addition of peroxydase reagent was measured spectrophotometrically, 25 detailed Materials and Methods. A comparative survey was made in with total parallel with A36 and (supernatant) of M. paratuberculosis (equal amounts of proteins were used for the two assays).

All the sera of infected animals (stages II and III of the Johne's disease) yielded a positive answer (values of 0.84 to 2.25 units) to both types of the ELISA assay (Figure 3). On the contrary, uninfected animals were invariably negative (values lower than 0.38 units). With A36-ELISA, considerably higher absorbance values (1.44 to 2.25 units) were obtained than with the total cytoplasm-ELISA (0.84 to 1.65).

These results suggest the immunodominance of the A36 antigen in the Johne's disease, and the usefulness of the A36-based ELISA as a diagnostic assay.

Peripheral location of the TMA complex in mycobacteria:

The observed immunodominance of A36 is more compatible with a surface component than with an antigen complex located in the cytoplasm. However, a transfer of TMA complex through the envelope and its protrusion at the cell surface is conceivable.

The use of the immunoelectron microscopy methodology has allowed a direct approach to this problem. Multiplying cells of M. paratuberculosis were incubated with anti-A36 Ig from immunized rabbits. Cell-bound primary antibodies were revealed by secondary swine anti-rabbit IgG labeled with colloidal gold. Electron micrographs show the presence of antigen reactive spots on the surface of mycobacteria (results not shown).

These data indicate that part of the TMA complex does indeed occur within the cell wall and is presented on the cell surface.

Immunological crossreactivity of A36 and other TMA antigens:

In the preceding section, the development of a A36-based ELISA assay for titration of antimycobacterial antibodies has been described. possible use of this assay in Veterinary Medicine relies on its specificity with respect to: a) other mycobacteria which are usual hosts of the intestinal and b) tracts of ruminants; M. bovis, tuberculosis which can cause tuberculosis in cattle (compulsory slaughtering of PPD-positive cattle). This problem evaluating asw approached by the crossreactivity of TMA complexes from different mycobacteria with two procedures (see Table 1).

A first series of assays was carried out with microtitration plates coated with the TMA complex from M. avium, M. bovis, M. paratuberculosis and M. phlei. All these plates were used to titrate a single anti-A36 antiserum, a procedure yielding an evaluation of the percentage of shared TMA epitopes. Considering the autologous reaction (A36-anti A36 IgG) equal to 100, percentage of homology of M. paratuberculosis TMA complex with the TMA complex of M. avium and bovis was very high; it was much lower for M. phlei TMA complex.

When the A36-based ELISA assay was repeated with anti-A36 antiserum previously absorbed by different mycobacterial TMA complexes, an evaluation of the A36 specific epitopes was obtained. From Table 1, it results that the percentage of specific epitopes was low when the A36 was compared to the TMA of M. avium and M. bovis, it was high when compared to the TMA of M. phlei.

FOUR OF COMPLEXES THA TABLE 1 : CROSSREACTING AND SPECIES SPECIFIC EPITOPES IN THE MYCOBACTERIA

THA In BLIBA

Parameter	Coating reagent (plate)*	Absorbing reagent (antiserum) ^b	ELISA unita (A _{450m}) ^c	Epitopes (%) Cross- specific ^d reacting
A. Crossreactivity	M. parat. M. bovis M. phlei	1111	2.367 2.376(±0.247) 2.240(±0.181) 1.083(±0.156)	100 100(±13) 96(±10) 49(± 8)
B. Specificity	M. parat. M. parat. M. parat.	H. parat. H. avium H. bovis H. phlei	0.462 0.574(±0.197) 0.603(±0.238) 1.073(±0.141)	0 7(±11) 10(±13) 48(± 8)

coat used Were μg/well) (0.5 mycobacteria different from microtitration plates preparations

b anti-A36 antiserum was pre-absorbed (samples B) or not (samples A) with TMA complex from different mycobacteria

ant1-A36 Ig were revealed by a second labeled (samples A) or with different TMAs to plates coated with A36 (samples B) or with antiserum (1/150000 dilution) was added, and bound antibody

^d percentage of crossreacting or specific epitopes calculated on a logarithmic scale.

These results show the lack of species-specificity of the A36-ELISA as a diagnostic reagent for the Johne's disease. They suggest, however, the possible occurrence of A36 components endowed with such a specificity.

Immunodominance and specificity of the A36 proteins:

The species specificity, which was missing at the level of the complete A36 antigen complex, was sought with respect to its proteins components. The TMA complexes from M. avium, M. bovis, M. paratuberculosis and M. phlei were isolated, and their fractionated by polyacrylamide components were electrophoresis. A similarity of M. avium and M. paratuberculosis tracks is apparent, whereas those of M. bovis and M. phlei TMA were clearly different to the M. paratuberculosis track.

When fractionated A36 proteins were immunoblotted with anti-A36 antiserum, a dozen of major polypeptides were stained, most of them located in the 28-42 kDa region. Immunoblotting with anti-A36 antiserum prelysate of M. phlei yielded absorbed with a polypeptide bands; they were 3 in the case of M. bovis and one with M. avium. Table 2 provides a comparative evaluation of the main A36 components according to two properties: immunogenicity level (staining intensity by and sera of infected bovines) pooled specificity (lack of cross-reactivity with the other mycobacteria). Eleven major components of 22 to 74 kDa are listed: two of them (of 23 and 31 kDa) containing specific epitopes with respect to the tested organisms except M. avium, and one of 34 kDa containing specific epitopes with respect to all of the tested organisms including M. avium.

OF COMPLEX. THA THE OF PROTEINS SOME **O**F CHARACTERISTICS TABLE 2 : IMMUNOLOGICAL CHARACTE MYCOBACTERIUM PARATUBERCULOSIS (A36)

		. 			_	-						
	M. phlei	ou	ou	уев	ou	уев	уев	уев	уев	уев	уев	уев
Spacificity ^d towards	M. bovis	ou	ou .	Ou	or C	no	pu	уев	уев	2	yes	OU .
	M. avium	ou -	on	on	ou	ou	ou	yes	2	ou	ou	ou
genicity ^{b,c} s in hosts)	Vines III	+	1		+	+	+	+++	+++	+	1	1
ty ^{b, c} hosts)	CTed bo	1	1	+	+	1	‡	+ + +		1	+	+
genicity ^{b,6} Is in host	T I	1	ı	+	+	‡	‡ ‡	+ + +	+++	1	1	1 .
Immunog (lavel	rabbit anti-A36	‡	+	+	+ + +	++	+	+ + +	‡	+ +	+++	+
Protein• (kDa)		74	52	41	.	37.	35	34	31	53	23	22

* A36 was dissociated and protein components were fractionated by SDS-PAGE electrophoresis and identified by immunoblotting

of the intensity b degree of immunogenicity for rabbits and cows was evaluated from immunoblot staining with the corresponding sera

c sera from cattle affected by different stages of the Johne's disease: I, asymptomaticnon excretory; II, asymptomatic-excretory; and III, symptomatic-excretory forms

crossreactivity was expressed by a no, and specificity by a yes.

The immunological relevance of the latter protein was checked by immunoblot analysis of A36 proteins with infected bovine sera: a major band at the level of the 34 kDa marker was observed (Figure 4, lanes 4, 5, 6 and 7). This band was missing in the control (lane 8 with healthy bovine serum).

It is thus evident that the 34 kDa protein component of the TMA complex is immunodominant in cattle, relevant to Johne's disease, and containing species-specific epitopes with respect to related mycobacteria.

The present invention enables to develop a A36 based ELISA test for paratuberculosis: its ability to reveal the presence of a mycobacterial infection in cattle has been proven in Figure 3. Basic requirements for the use of a given antigen as reagent medical interest of are: 1) its immunoassays its relevance to the targeted immunodominance; 2) disease; and 3) its specificity. Requirements 1 and 2 were therefore fulfilled by the A36 based-ELISA. Requirements 1 to 3 are completely fulfilled by the p362 polypeptide which is part of the 34 kDa protein belonging to A36, as described hereafter.

EXAMPLE II: Isolation of clone a362 expressing a 10 kDa polypeptide (p362), DNA sequencing of the insert of clone a362 and testing of p362 in an ELISA for Johne's disease:

MATERIAL AND METHODS

Cloning vectors

The following types were used: Agt11 (Young R.A. and Davis R.W., 1983, "Yeast RNA polymerase II genes: isolation with antibody probes" Proc. Natl. Acad. Sci. USA 80:1195-1198) and pUEX2 (Brennan G.M. et al., 1987, "pUEX, a bacterial expression vector related to pEX with universal host specificity" Nucl. Acids Res.

15:10056) and pmTNF-MPH (see Figures 9a, 9b and Table 5) as expression vectors, and the Blue-Script SK as sequencing vector (Stratagene).

Bacteria

Mycobacterium paratuberculosis 19698 (from the American Type Culture Collection). M. paratuberculosis: strain 2887 (Crohn): ATCC n° 43015. M. avium serotype 4, M. avium serotype 2, M. avium serotype 8 (Schaefer 1965, "Serologic identification classification of the atypical mycobacteria by their agglutination" Am. Rev. Resp. Dis. suppl. 92:85-93). M. tuberculosis H37rv: ATCC n° 25618. M. gordonae: ATCC n° 14470. Brucella abortus B3 (Cloeckaent A. et al., 1990, Infect. Immun. 58:3980-3987). Strains of Escherichia coli: Y1089 (Δ(lacU169), Δ(lon), hflA150 (chr::Tn10), (pMC9), (rK, mK)), Y1090 ($\Delta(\frac{1ac}{0169})$, $\Delta(\frac{1on}{0})$, sup F, $(trpC22::Tn10), (pMC9), (rK, mK)), MC1061 (<math>\Delta(lacX74)$, galU, galK, (rK, mK')) and DH5aF' (F', (rK, mK'), supE44, lacZAM15, A(lacZYA argF) U169), K12AH, ATCC 33767 (lacZ(am) Δ (bio uvr B) (λ Nam7 am53 cI 857 Δ H1) rpsL20).

Antisera

Rabbit anti-M. paratuberculosis antiserum was from Dako (Copenhagen, Denmark, lot n° 014). Sera from paratuberculosis-infected cattle were provided by Dr. M. Desmecht (National Institute for Veterinary Research, Brussels) and Dr. B. Limbourg (Erpent, Center of Veterinary Medicine, Belgium).

Polyclonal antisera against whole homogenate of $\underline{\text{M.}}$ avium serotype 4, $\underline{\text{M.}}$ bovis BCG, and $\underline{\text{M.}}$ phlei, as well as those against the TMA complex and β gal-p362 (recombinant polypeptide of the invention fused to β -galactosidase hereafter described) were produced by repeated subcutaneous inoculations into rabbits (10 μ g proteins/0.5 ml buffered saline emulsified with equal

volume of incomplete Freund's adjuvant, 6 inoculations at 1-week intervals).

Purification of M. paratuberculosis DNA:

Suspensions of acteria (10 mg in 0.5 ml of 100 mM NaCl, 1 mM EDTA, 50 MM Tris-HCl pH 7,4) were incubated sequentially with lysozyme (25 μ l of 20 mg/ml, 14 h, 50°C), pronase (25 μ l of 20 mg/ml, 1 h, 37°C), and SDS (25 μ l of 20%, 1 h 37°C). Mixtures were extracted with chloroform-isoamyl alcohol (24:1, vol:vol), watersaturated phenol, and ether. After incubation with ribonuclease (5 μ l of 2 mg/ml, 1 h, 37°C), DNA was purified on columns of Sephadex G50 (equilibrated with 4,8 mM sodium phosphate pH 6,8) and hydroxyapatite (washed with 8 M urea, 0,1 M sodium phosphate buffer pH 6,8 containing 1% SDS, and then with 4,8 mM sodium phosphate pH 6,8, and eluted with 480 mM sodium phosphate pH 6,8).

Construction of a lqt11 library of M. paratuberculosis:

M. paratuberculosis DNA was sheared to average length segments of 0,5 to 1,5 kb ultrasonicator 60 W, 2 sec). Shearing was monitored by (Vibra Cell electrophoresis. gel EcoR1 sites methylated with EcoRl methylase (5 μg of sheared DNA in 50 μ l of buffer (50 mM Tris-HCl pH 7,5, 1 mM Na₃EDTA, 5 mM dithiothreitol, 50 μ M S-adenosyl-L-methionine and 10 units of EcoR1 methylase). Methylation was pursued for 30 min at 37°C, and stopped by 10 min incubation at 70°C. Blunt-end DNA fragments were incubation with T4 DNA polymerase (5 μ l of 0,1 M MgCl₂, obtained 2,5 μ l of 1 mM dTNPs, 1 μ l of 1 M(NH₄)₂SO₄, and 20 units of T4 DNA polymerase per 40 μ l methylation reaction medium; 20 min incubation at 37°C). EDTA (15 mM final concentration) Was added, reaction mixture extracted with phenol/chloroform twice, and the aqueous phase was extracted with ether. After addition of sodium acetate 0,3 M final concentration, DNA was

precipitated with 2 vol of EtOH at -20°C and washed with 70% EtOH. DNA pellet was dissolved in buffer (10 μ l of 100 mM Tris-HCl pH 7,5, 20 mM MgCl₂, 20 mM dithiothreitol), phosphorylated EcoR1 linkers μ g/ml) were added, followed by addition of PEG 6000 (final concentration 15%), 1 mM ATP concentration) and 2 units of T4 DNA ligase, and the reaction mixture was incubated overnight at 12°C. This mixture was incubated at 37°C with an excess of EcoRl, and DNA fragments were purified from linker excess on Sephadex G25. The DNA solution thus obtained was extracted sequentially with phenol/chloroform and ether, precipitated, and washed with ethanol. pellet (0,5 μ g) was dissolved in TE buffer (10 mM Tris-HCl pH 7,5, 0,1 mM EDTA) and ligated (18 h, 4°C) with 1 µg of dephosphorylated EcoR1-digested Agt11 DNA (Promega). Methylation, ligation, and digestion steps were controlled by agarose gel electrophoresis. Phage packaging of cloned DNA was obtained with Stratagene gigapack extract.

Screening of the Agt11 library and dot-blot technique:

After infection of <u>E. coli</u> Y1090 by the recombinant phage mixture and spreading them out over the plate, they were incubated for 3-4 h at 42°C.

For identification of recombinant phages, IPTG (isopropylthio β-galactopyranoside) (10 mM) saturated nitrocellulose filters were placed directly on the surface of the overlay plates containing the plaques and incubated for 18 h at 37°C (Young R.A. and Davis R.W., 1983, "Yeast RNA polymerase II genes: isolation with antibody probes" Proc. Natl. Acad. Sci. USA 80:1195-1198). After spotting of control antigens (1 μg) and washing for 10 min with TBS buffer (0,5 M NaCl, 0,023 M Tris-HCl pH 7,5), filters were incubated for 30 min with the same buffer containing 3% (w/v) gelatin and then with the rabbit anti-M. paratuberculosis

antiserum (Dako) previously diluted with TBST buffer (TBS buffer containing 0,05% (v/v) Tween 20) containing 1% (w/v) gelatin. After washing, filters were incubated for 1 h with 1/400 dilutions of peroxydase-labeled anti-rabbit Ig. After repeated washing with TBST and TBS, the peroxydase substrate α -chloronaphtol (Bio Rad Laboratories, Richmond, Calif.) and hydrogen peroxide were added. Reaction was stopped by washing with distilled water. Plaques corresponding to reactive spots on the filters were picked off, transferred to SM medium (100 mM NaCl, 10 mM MgSO,, 20 mM Tris-HCl pH 7,4) and purified by repeated passages in E. coli Recombinant clones were then further their characterized with respect to antigenicity (incubation with bovine sera and anti-A36) and their specificity (incubation with antibodies against homogenate of M. avium, M. bovis and M. phlei) using the same procedure as described above.

A similar technique was used for dot-blot experiments in which the specificity of the recombinant polypeptide p362 was tested with respect to different mycobacteria: spots of mycobacterial homogenates on nitrocellulose membranes were incubated with anti- β gal-p362 Ig.

High level expression of fusion protein in E. coli:

Colonies of <u>E. coli</u> Y1089 lysogenized with the appropriate λ gtll recombinants were multiplied at 30°C in Luria-Bertani medium ($A_{600\text{rm}}$ =0,5). After heat shock (20 min at 45°C), production of β -galactosidase fusion proteins of the invention was induced by the addition of 10 mM IPTG (final concentration) and further incubation (60 min at 37°C). Cells harvested by centrifugation were suspended in buffer (10 mM Tris-HCl, pH 8,2, 2 mM EDTA) and rapidly frozen in liquid nitrogen.

For enhanced expression, Agt11 inserts were subcloned into the expression vector pUEX2 (Brennan G.M. et al., 1987, "pUEX, a bacterial expression vector related to pEX with universal host specificity" Nucl. Acids Res. 15:10056), commercially available from Amersham, which was used to transform E. coli MC1061 (Maniatis, Molecular Cloning). Single colonies of transformed E. coli were grown at 30°C to A₆₀₀=0,3 and heat-shocked (90 min at 42°C). Harvested cells were lysed by sonication and frozen in liquid nitrogen. Protein fractionation and immunoblotting:

The TMA complex and recombinant proteins were analyzed by polyacrylamide gel electrophoresis under

denaturing conditions (SDS PAGE) (Laemmli, U.K. 1970, "Cleavage of structural proteins during the assembly of

the head of bacteriophage T4", Nature 227:680-695).

Fractionation on 7,5 or 10% acrylamide gels was carried out in a 2001 vertical electrophoresis unit (LKB-Produkter AB, Bromma, Sweden) (4 h, 50 V, 20°C). Molecular weight protein markers (Sigma, St Louis, Mo) were: myosin (205 kDa), β -galactosidase (116 kDa), phosphorylase B (97,4 kDa), bovine serum albumin (66 kDa), ovalbumin (45 kDa), glyceraldehyde-3-phosphate dehydrogenase (36 kDa) carbonic anhydrase (29 kDa), trypsinogen (24 kDa), trypsin inhibitor (20.1 kDa), and a-lactalbumin (14.2 kDa). Protein bands were stained with Coomassie brilliant blue. Electrophoresed proteins were transblotted (LKB 217 Multiphor 2 Electrophoresis System, 10 V, 2 h, with buffer 20% methanol, 0,039 M glycine and 0,048 M Tris base, pH 8.8) nitrocellulose membranes. Mycobacterial antigens were visualized by sequential incubation with polyclonal rabbit antisera (anti-A36 for recombinant mycobacterial antigens fused to β -galactosidase or anti- β gal-p362 for TMA proteins) and peroxydase-labeled anti-rabbit Ig (Dako, Copenhagen, Denmark) (1/400 dilution). Total

protein blotted on the membrane was visualized by staining with India ink.

DNA Sequencing:

Sequence analysis of the DNA insert of recombinant clone a362 was done by the primer extension and dideoxy termination method (Sanger F. et al., 1977, "DNA sequencing with chain terminating inhibitors", Proc. Natl. Acad. Sci. USA 74:5463-5467), subcloning of the Agtll insert into the sequencing pBluescript SK (Stratagene). vector Sequencing reactions were performed with T7 DNA polymerase and different primers (universal, reverse, SK, primers from Deaza Kit, Pharmacia, Uppsala, Sweden). Computer-aided analysis of nucleic acid and polypeptide sequences were performed with the program COD-FICK (PC-GENE, Intelligenetics, USA). Homology searches were performed on DNA level in EMBL bank (release 29) and UGEN bank (release 70-29) (Intelligenetics Inc., CA-USA), and on protein level in PIR bank (release 31) and Swiss Prot (release 20). No homologous sequences were found.

Serological analysis (ELISA) with recombinant polypeptides:

Multiwell microtiter plates (Microwell Module, High binding Capacity, Nunc, Denmark) were coated with total cytoplasm of <u>E. coli</u>—a362 and with total cytoplasm of <u>E. coli</u> as a control. Four μ g of soluble proteins / 50 μ l 0,05 M Na carbonate buffer pH 9,6 were coated per well. Plates were air dried overnight and saturated (0,1% serum albumin in 0,15 M NaCl, 1 h at 37°C). Dilutions of bovine Ig in PBST (0,15 M NaCl, 0,02 M phosphate buffer pH 7,2, containing 0,005% Tween 80) were added to plate wells (50 :1, 1 h at 37°C). Peroxydase—labelled rabbit anti-cow Ig (Dako) (50 μ l, 1/400 dilution in PBST/per well) were added (1 h at 37°C). Excess of reagent was removed by 5 PBST

washings. After incubation with peroxydase reagent (50 μ1/well of 0.2% 0-phenylenediamine with 0,015% hydrogen peroxyde in 0,017 M Na citrate buffer pH 6,3, 30 min, 37°C in the dark), the reaction was stopped with 50 μ l 2 M H₂SO₄, and A_{450rm} was measured in a colorimetric plate reader (SLT 210, Kontron Analytical, UK). Results were recordered as ELISA absorbance values. In some experiments, CTOSS reactive Ig were incubation (18 h at 4°C) with bacterial homogenate. Absorbed preparations were checked by dot-blot trials before applications in immunoblots or immunoassays. Immune electron microscopy:

Suspensions of mycobacteria in water (5×10^7) cells/5 μ l) were placed on carbon-formvar 200-mesh copper grids and air-dried. Grids were serially incubated with: a) bovine serum albumin (3% solution in buffered saline, 30 min, 37°C); b) anti-\betagal-p362 rabbit antiserum (a 10.3 dilution of Ig in buffered saline with 0,05% Tween 20, 2 h, 37°C); c) sheep antirabbit biotinylated Ig (1/200 dilution of Ig from Amersham, U.K., in buffered saline-Tween, 1 h, 20°C); d) gold-labelled streptavidin (a 1/20 dilution of a preparation from Amersham, U.K.) (Cloeckaert A. et al., 1990, "Identification of seven surface-exposed Brucella outer membrane proteins by use of monoclonal antibodies: immunogold labeling for electron microscopy and enzyme-linked immunosorbent assay", Infec. Immun. 58:3980-3987).

Grids were analyzed in a transmission electron microscope (Philips CM 10).

RESULTS

1. Preparation of a genomic library of M. paratuberculosis and isolation of recombinant clones:

A genomic library of <u>M. paratuberculosis</u> has been prepared by the use of the expression vector \(\alpha \text{gtll.} \) For this purpose, purified mycobacterial DNA was sonicated

under controlled conditions yielding segments of 103 bp on the average (0.5 to 2×10^3). These fragments were methylated by EcoR1 DNA methylase (efficiency of methylation was controlled by incubation with EcoR1), incubated with T4 DNA polymerase to obtain blunt-end DNA, and provided with EcoR1 linkers by incubation with T4 DNA ligase. After EcoR1 digestion, DNA segments were purified free of linker excess and inserted into EcoR1-cleaved Agt11 by incubation with T4 DNA ligase (a step checked by gel electrophoresis). After packaging and infection of E. coli Y1090, 7.5 x 105 recombinant clones (75% of total clones) were obtained, one third screened with rabbit of which was anti-M. paratuberculosis antiserum (Dako). After repeated purifications, ten recombinant clones were selected: three of them expressed TMA complex proteins, and seven produced epitopes of proteins not present within the TMA complex.

2. Analysis of antigenicity and specificity of polypeptides produced by recombinant clones:

Since cloning of <u>M. paratuberculosis</u> genes was aimed at producing polypeptides to be used as diagnostic reagents, it appeared essential to test the reactivity of recombinant clones towards sera of cattle affected by the Johne's disease. As shown in Table 3, all the selected clones reacted with sera of animals bearing one of the clinical forms of the disease. The strongest reactions were afforded by clones a4 and a362. On the contrary, no reactivity was observed with sera from healthy bovines.

TABLE 3
Characteristics of clones expressing an antigenic polypeptide of <u>M. paratuberculosis</u>

Clones*	Ant	igeni	city**	Specificity with respect to						
	1	2	3	M. avium	M. bovis	M. phlei				
										
a1	(+)	+	+.	no	no	yes				
a2 '	+	+	+	yes	yes	yes				
a3	+	+	++	no	yes	yes				
a4	++	++	++	no	no	yes				
a 5	+	+	+ ·	no	yes	yes				
a 6	+	+	++	no	no	yes				
a7	(+)	+	+	ָסת	no	no				
a361	+ .	+	++	no	yes	yes				
a362	++	++	++	yes	Yes	yes				
a363	(+)	+	+	no	no '	yes				

^{*} only clones a361 to a363 express polypeptides belonging to the TMA complex.

Another requirement of paramount importance was the specificity with respect to mycobacteria belonging to the saprophytic and pathogenic flora of cattle. Recombinant clones were tested for reactivity with

^{**} detected by sera from asymptomatic and non
excretory bovine (1), asymptomatic and excretory
bovine (2) and symptomatic and excretory bovine
(3); quantified as low reaction "(+)", good
reaction "+" and very good reaction "++".

^{***} cross reactivity was expressed by a "no", and specificity by a "yes".

antisera against homogenates of <u>M. avium</u>, <u>M. bovis</u> and <u>M. phlei</u>. It was previously shown that the overall DNA homology levels of these three mycobacteria with respect to <u>M. paratuberculosis</u> were respectively 94, 52, and 19 percent (Hurley S.S. et al., 1988, "DNA relatedness of <u>M. paratuberculosis</u> to other members of the family of mycobacteriaceae", Int. Journal Syst. Bact. <u>38</u>:143-146). Data in Table 3 indicate that, although all clones but one were specific towards <u>M. phlei</u>, only five of them were specific for <u>M. bovis</u> and two for <u>M. avium</u>.

In conclusion, only two of the selected clones, a2 and a362 fulfilled both requirements for species-specificity and relevance to Johne's disease. Moreover, only the latter clone reacted with anti-A36 antiserum and corresponded, therefore, to a A36 protein, presumably the 34 kDa protein previously identified as a TMA complex component with species-specific epitopes. The remaining part of this example relates to the characterization and use of clone a362.

3. Size of clone a362 insert and its expressed polypeptide p362:

EcoR1 cleavage of DNA of clone a362 yielded an insert of about 500 bp devoid of internal EcoR1 restriction sites (not shown).

E. coli Y1089 was lysogenized by the recombinant phage, and the synthesis of a chimaeric protein fused with β -galactosidase was induced by IPTG: a fusion protein of about 125 kDa (β gal-p362) was produced (Figure 5). Since β -galactosidase (116 kDa) misses 2 kDa in λ gtll, the recombinant polypeptide coded for by the insert of clone a362 (p362) is expected to be about 11 kDa in size. Consequently, only a roughly 300 bp portion of the 500 bp insert coded for such an 11 kDa polypeptide. This was confirmed by sequencing and

determination of the orientation of the insert DNA as described further.

4. Production of p362 recombinant polypeptide and evidence of its belonging to a 34 kDa protein of A36:

Since the production of the β -gal p362 by <u>E. coli</u> Y1089 containing the λ gtll-recombinant phage was only 2% of total proteins, the corresponding insert was recloned in a more favorable expression vector. For this purpose, the λ gtll insert of the a362 recombinant clone was freed by incubation with EcoRl, purified by electroelution from an agarose gel (75% recovery), and recloned into the EcoRl site of the expression vector pUEX2 (Amersham). In this case, production of β gal-p362 fusion protein in the transformed MC1061 strain of <u>E. coli</u> (6 x 10⁵ transformants/ μ g DNA) was about 25% of total proteins.

After running the SDS-PAGE of the lysate from the transformed strains, the recombinant fusion protein was eluted from the polyacrylamide gel and used to elicit antibodies in rabbits (anti- β gal-p362).

The protein components of the TMA complex from M.

paratuberculosis were fractionated by electrophoresis
on polyacrylamide gels (SDS PAGE). After transfer to
nitrocellulose sheets, TMA proteins were incubated with
anti-\(\beta\gai\) and page 6, a major band
corresponding to the 34 kDa protein of the TMA complex
was immunolabeled: this was the unique TMA protein
containing species-specific epitopes as above reported.
A second band of about 31 kDa was stained to minor
extent: it was also present in the immunoblots of TMA
proteins with sera of infected cattle.

5. Localization of the p362 polypeptide at the bacterial surface:

Since the A36 antigen complex was previously shown to be present at the cell surface, a peripheral location of the p362 recombinant polypeptide would

further confirm the belonging of p362 recombinant polypeptide to a protein of the A36 complex. Electron micrographs show indeed the presence of the p362 polypeptide within the cell wall and its release during the declining growth phase (results not shown).

6. Assessment of the species-specificity of the recombinant polypeptide p362:

From what is above-mentioned, it is shown that the 34 kDa protein component of the TMA complex of M. paratuberculosis contains epitopes devoid crossreactivity towards M. bovis, M. avium and M. phlei. Although the recombinant p362 polypeptide, which apparently represents a portion of the 34 kDa protein, is likely to be endowed of species-specificity, a more stringent confirmation is needed for a polypeptide reagent forecast for serological the specificity of p362 was tested Consequently, against two series of M. paratuberculosis and M. avium isolates from cattle as well as against certain Grampositive and Gram-negative bacteria being usual hosts of bovine gut (Table 4).

The dot-blot experiment was carried out by spotting on a nitrocellulose membrane 2 μ g samples of different bacterial homogenates. Membranes were then incubated successively with rabbit anti- β gal-p362 antiserum and, after washing, with peroxydase-labeled swine anti-rabbit IgG. Spots were revealed by the peroxydase reaction. All of eight \underline{M} . paratuberculosis isolates were positive, whereas the closely related organisms of the MAIS group were negative. None of the other tested mycobacteria gave a positive reaction, neither did the Nocardia and Brucella species (see Table 4).

[HYCO]BACTERIA
OTHER
62 TOWARDS OTI
p 362
90
SPECIFICITY OF p362 TOWARDS OTHER [MYCO]BAG
4
TABLE

Bacterium lysates	Anti-Agal-p362	Bacterium lysates	Ant1-βgal-p362
- M. paratuberculosis: 2E 316F ATC 19698 ATC 43015 2890(bovine)(1) 2891(bovine)(1) 2895(goat)(1) 172 28/66(bovine)(2) - M. avium D4(5) - M. avium serotype 8 - M. avium serotype 8 - M. avium serotype 9 - M. avium serotype 1 - M. avium serotype 1 - M. avium serotype 2 - M. avium serotype 2 - M. scrofulaceum(1) - Salmonella typhimurium(3)	+++++++11111	M. intracellulare (1) MAIS A3(4) MAIS A84(4) MAIS 8715(4) MAIS 87537(4) M. bovis BCG GL2 M. tuberculosis H37rv(6) M. phlei AM76(1) M. ieprae(1) M. ieprae(1) M. gordonae ATCC 14430 Nocardia asteroides(1) Brucella abortus B3(3)	

(-) absence of reaction positive immunological reaction £325409

Portaels IMTA (Institut de Médecine Tropicale, Anvers Belgique) from Kaeckenbeeck DBUL (Département de Bactériologie, Université de Liège, Belgique) from LIMET ICP (Institut of Cellular Pathology, Belgique) from Defoe IPB (Institut Pasteur du Brabant, Belgique) from Saxegaard NVIN (National Veterinary Institute, Norway).

25618 ATCC

7. Sequencing of the cloned insert coding for polypeptide p362:

To sequence the 500 bp DNA segment coding for the polypeptide p362, the insert of clone a362 was isolated by EcoR1 cleavage from the chimaeric vector λ gtl1 and recloned into the Bluescript vector SK. After transformation of <u>E. coli</u> DH5 α F¹, clones carrying inserts coding for p362 were selected.

The sequence of the insert showed the occurrence 507 bp DNA segment flanked by two EcoR1 extremities (Figure 7C). The G+C content of this segment was 70%, in agreement with the 64% G+C of the whole M. paratuberculosis genome. The sequence in Figure 7C yielded two open reading frames in phase with the EcoRI sites: a 306 bp region (1 to 306) in one direction, and a 185 bp region (507 to 322) into opposite orientation. The program COD-FICK (PC-GENE) which takes in account the codon usage, confirmed the coding ability of the two open reading frames. They coded respectively for 10 kDa and 7 kDa polypeptides. The insert was subcloned in an expression vector in E. coli in both orientations. Only one orientation yielded an expression product reacting with the rabbit antißgal-p362 antiserum. Restriction analysis led to the selection of the 306 bp open reading frame as being the one coding for the p362 polypeptide [10 kDa]. selected coding region and the aminoacid sequence of polypeptide p362, corresponding to the carboxyterminal extremity of the 34 kDa protein are displayed in Figure 8.

8. Testing of p362 in an ELISA for Johne's disease:

The 10 kDa polypeptide (p362), endowed with species-specificity, and being part of the 34 kDa protein of A36, can be used as a specific test for paratuberculosis.

A preliminary test has been done using plates coated with total cytoplasm of <u>E. coli</u>-a362 containing p362. Bovine sera were preabsorbed to <u>E. coli</u>-control homogenate. Figure 2 shows that all sera from infected bovines react significantly with p362. On the contrary, healthy bovines (samples 26-32) do not give a signal which is significantly higher than that observed with <u>E. coli</u>-control cytoplasm.

Antibodies directed against p362 are already present in the early stages of the disease (samples 1-13). p362 can thus be considered as a very suitable antigen for specific and sensitive diagnosis of paratuberculosis.

To decrease the background levels due to cross reaction with the β -galactosidase part of the fusion protein, the insert coding for p362 was recloned into another: expression vector (pmTNF-MPH, Innogenetics) (Figures 9a and 9b).

It contains the tetracycline resistance gene and the origin of replication of pATiss (obtainable from Bioexcellence, Biores B.V., Woerden. The Netherlands), the lambda PL promoter up to the MboII site in the N gene 5' untranslated region (originating from pPL(λ); Pharmacia), followed by a synthetic ribosome binding site (see sequence data), and the information encoding the first 25 AA of mTNF (except for the initial Leu which is converted to Val). This sequence is, in turn, followed by a synthetic polylinker sequence which encodes six consecutive histidines followed by several proteolytic sites (a formic acid, CNBr, kallikrein, and E. coli protease VII sensitive site, respectively), each accessible via a different restriction enzyme which is unique for the plasmid (SmaI, NcoI, BspMII and Stul, respectively; see restriction and genetic map, Figure 9a). Downstream from the polylinker, several transcription terminators are present including the E.

<u>coli</u> trp terminator (synthetic) and the $rrnBT_1T_2$ (originating from pKK223-3; Pharmacia). The total nucleic acid sequence of this plasmid is represented in Figure 9b.

Table 5 gives a complete restriction site analysis of pmTNF-MPH.

The presence of 6 successive histidines allows purification of the fusion protein by Immobilized Metal Ion Affinity Chromatography (IMAC).

To subclone the insert coding for p362 in pmTNF-MPH, it was set free from the construct in vector pUEX2 by EcoRI digestion. The EcoRI fragment (507 bp) was eluted from the gel, purified, blunted and inserted in the blunted XbaI site of pmTNF-MPH. The resulting recombinant plasmid, pmTNF-MPH-a362, is brought into E. coli strain Kl2AH (ATCC 33767) by transformation. After growth at 28°C, expression of the recombinant protein is induced by a temperature shift to 42°C, which is held on during 2 hours. Cells were harvested, centrifuged and lysed in French press.

The expressed fusion protein mTNF-H6-p362, present in the cytoplasm fraction of the <u>E. coli</u> recombinant, is purified by Immobilized Metal Ion Affinity Chromatography (IMAC) using conditions known by the man skilled in the art. The amino acid sequence of this complete fusion protein is represented in Figure 10.

The purified fusion protein is used to coat 96-well microtitration plates, which were incubated with serial dilutions of sera from uninfected (control) and infected animals. Plate bound IgG were titrated with peroxydase-labeled rabbit anti-bovine IgG, as described in Materials and Methods.

CLAIMS

- 1. Polypeptide containing in its polypeptidic chain:
- the amino acid sequence of 101 amino acids of Figure 8,
- or a fragment of this sequence, this fragment being such that:
 - . it is liable to be recognized by antibodies also recognizing the abovesaid sequence of 101 amino acids, but it is not recognized by antibodies respectively raised against M. bovis, M. avium, M. phlei and M. tuberculosis, and possibly against M. leprae, M. intracellulare, M. scrofulaceum, M. fortuitum, M. gordonae and M. smeqmatis,
 - . it is liable to generate antibodies which also recognize the abovesaid sequence of 101 amino acids but which do not recognize M. bovis, M. avium; M. phlei and M. tuberculosis, and possibly M. leprae, M. intracellulare, M. scrofulaceum, M. fortuitum, M. gordonae and M. smegmatis,
 - . it reacts with the majority of sera from cattle suffering from Johne's disease,
- or the polypeptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the above-mentioned properties.
- 2. Polypeptide according to Claim 1, characterized by the fact that it is constituted by the sequence represented on Figure 8, extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (101), or by the following peptides:

Glu-Phe-Pro-Gly-Gly-Gln-Gln-His-Ser-Pro-Gln,

Gln-Gln-Ser-Tyr-Gly-Gln-Glu-Pro-Ser-Ser-Pro-Ser-Gly-Pro-Thr-Pro-Ala.

- 3. Polypeptide according to Claim 1, characterized by the fact that:
- it contains the amino sequence of 101 amino acids of Figure 8 as its C-terminal part,
- it has a molecular weight of about 34kDa, in SDS-PAGE,
- it is coded by a nucleotide sequence liable to hybridize with the complementary strand of the sequence of Figure 11,
- it reacts with the majority of sera from cattle suffering from Johne's disease,
- it is advantageously liable to elicit a cellular immune response in sensitized subjects.
- 4. Amino acid sequences constituted by anyone of the polypeptides according to Claims 1 to 3 and a protein or an heterologous sequence with respect to said polypeptide, said protein or heterologous sequence comprising for instance from about 1 to about 1100 amino acids.
- 5. Nucleic acid characterized by the fact that it comprises or is constituted by:
- a nucleotide chain liable to hybridize with the nucleotide chain coding for the polypeptides according to anyone of Claims 1 to 3, or
- a nucleotide chain coding for the polypeptides
 according to anyone of Claims 1 to 3, or
- the complementary sequences of the above nucleotide chains.
- 6. Nucleic acid according to Claim 5, characterized by the fact that it comprises or is constituted by a nucleotide chain,
- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (307) on Figure 7A, or

or

- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (508) on Figure 7A, wherein
- X and E represent phosphodiester bonds, Y and F represent respectively G and C, Z and H represent respectively C and G,
- X and E represent respectively G and C, Y and F represent respectively C and G, Z and H represent phosphodiester bonds.
- 7. Nucleic acid according to Claim 5, characterized by the fact that it comprises or is constituted by a nucleotide chain,
- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (306) on Figure 7C, or
- extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (507) on Figure 7C.
- 8. Nucleic acid according to Claim 5, which comprises or is constituted by:
- a nucleotide sequence liable to hybridize with the complementary strand of the nucleotide sequence of Figure 11, or with the complementary strand of the nucleotide sequence extending from nucleotide at position (742) to nucleotide at position (1338) of Figure 11,
- the nucleotide sequence of Figure 11" or the nucleotide sequence extending from nucleotide at position (742) to nucleotide at position (1338) of Figure 11,
- the complementary sequences to the above-defined sequences.
- 9. Recombinant nucleic acid containing at least one of the nucleotide sequences of anyone of Claims 5

to 8 combined to or inserted in a heterologous nucleic acid.

- 10. Recombinant vector particularly for cloning and/or expression, comprising a vector sequence, notably of the type plasmid, cosmid, phage or virus DNA and a recombinant nucleic acid according to anyone of Claims 5 to 8, inserted in one of the non essential sites for its replication.
- 11. Recombinant vector according to Claim 10, containing necessary elements to promote the expression in a cellular host of polypeptides coded by nucleic acids according to anyone of Claims 5 to 8 inserted in said vector and notably a promoter recognized by the RNA polymerase of the cellular host, particularly an inducible promoter and possibly a sequence coding for transcription termination and possibly a signal sequence and/or an anchoring sequence.
- 12. Recombinant vector according to Claim 10, containing the elements enabling the expression by \underline{E} . coli of a fusion protein consisting of the polypeptide of β -galactosidase or part thereof linked to a polypeptide coded by a nucleic acid according to anyone of Claims 5 or 8.
- 13. Cellular host chosen from among bacteria such as <u>E. coli</u> or chosen from among eukaryotic organisms, such as CHO cells or insect cells, which is transformed by a recombinant vector according to anyone of Claims 9 to 12, and containing the regulation elements enabling the expression of the nucleotide sequence coding for the polypeptide according to anyone of Claims 1 to 3 in this host.
- 14. Expression product of a nucleic acid expressed by a transformed cellular host according to Claim 13.
- 15. Antibody characterized by the fact that it is specifically directed against a polypeptide according to anyone of Claims 1 to 3, and preferably by the fact

that it recognizes neither M. bovis, nor M. avium, nor M. phlei, nor M. tuberculosis.

- 16. Process for preparing a recombinant polypeptide according to anyone of Claims 1 to 4 comprising the following steps:
- the culture in an appropriate medium of a cellular host which has previously been transformed by an appropriate vector containing a nucleic acid according to anyone of Claims 5 to 8, and
- the recovery of the polypeptide produced by the above said transformed cellular host from the above said culture medium or from the cellular host.
- 17. Method for the <u>in vitro</u> diagnosis of paratuberculosis in an animal liable to be infected by <u>Mycobacterium paratuberculosis</u> comprising
- contacting a biological sample taken from an animal with a polypeptide according to anyone of Claims 1 to 3, or the expression product according to Claim 14, under conditions enabling an in vitro immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and the in vitro detection of the antigen/antibody complex which has been possibly formed.
- 18. Method for the <u>in vitro</u> diagnosis of paratuberculosis in an animal liable to be infected by <u>M. paratuberculosis</u>, comprising the following steps:
- contacting a biological sample with an appropriate antibody according to Claim 15, under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. paratuberculosis</u> which are possibly present in the biological sample and the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.
 - 19. Method for the <u>in vitro</u> diagnosis of Crohn's disease in a patient liable to be infected by

Mycobacterium paratuberculosis comprising the following steps:

- contacting a biological sample with an appropriate antibody according to Claim 15, under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. paratuberculosis</u> which are possibly present in the biological sample and the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.
- 20. Method for the <u>in vitro</u> diagnosis of Crohn's disease in a patient liable to be infected by <u>M. paratuberculosis</u>, comprising the following steps:
- contacting a biological sample taken from a patient with a polypeptide according to anyone of Claims 1 to 3, or the expression product according to Claim 14, under conditions enabling an in vitro immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and the in vitro detection of the antigen/antibody complex which has been possibly formed.
- 21. Necessary or kit for an in vitro diagnosis method of paratuberculosis in an animal liable to be infected by Mycobacterium paratuberculosis according to Claim 17, comprising:
- a polypeptide according to anyone of Claims 1 to 3, or the expression product of Claim 14,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned polypeptide is not labeled.
- 22. Necessary or kit for an in vitro diagnosis method of paratuberculosis in an animal liable to be

infected by <u>Mycobacterium paratuberculosis</u> according to Claim 18, comprising:

- an antibody according to Claim 15,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction, said reagents possibly having a label or being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned antibody is not labeled.
- 23. Necessary or kit for an in vitro diagnosis method of Crohn's disease in a patient liable to be infected by Mycobacterium paratuberculosis according to Claim 19, comprising:
- an antibody according to Claim 15,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction, said reagents possibly having a label or being liable to be recognized by a label reagent, more particularly in the case where the above mentioned antibody is not labeled.
- 24. Necessary or kit for an <u>in vitro</u> diagnosis method of Crohn's disease in a patient liable to be infected by <u>Mycobacterium paratuberculosis</u> according to Claim 20, comprising:
- a polypeptide according to anyone of Claims 1 to 3, or the expression product of Claim 14,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or being liable to be recognized by a labeled reagent,

more particularly in the case where the above mentioned polypeptide is not labeled.

- 25. Immunogenic composition comprising a polypeptide according to anyone of Claims 1 to 3, or the expression product of Claim 14, in association with a pharmaceutically acceptable vehicle.
- 26. Vaccine composition comprising among other immunogenic principles anyone of the polypeptides according to anyone of Claims 1 to 3 or the expression product of Claim 14, possibly coupled to a natural protein or to a synthetic polypeptide having a sufficient molecular weight so that the conjugate is able to induce in vivo the production of antibodies neutralizing Mycobacterium paratuberculosis, or induce in vivo a protective cellular immune response by activating M. paratuberculosis antigen-responsive T cells.
- 27. Necessary or kit for the diagnosis of prior exposure of an animal to <u>M. paratuberculosis</u>, said necessary or kit containing a preparation of at least one of the polypeptides or peptides according to anyone of Claims 1 to 3, or the expression product of Claim 14, with said preparation being able to induce <u>in vivo</u> after being intradermally injected to an animal a delayed type hypersensitivity reaction, at the site of injection, in case the animal has had prior exposure to <u>M. paratuberculosis</u>.
- 28. Polypeptides according to claim 3, characterized in that they contain or are constituted by:
- the amino acid sequence of Figure 11 or
- the amino acid sequence extending from amino acid at position (1) to amino acid at position (199) of Figure 11.

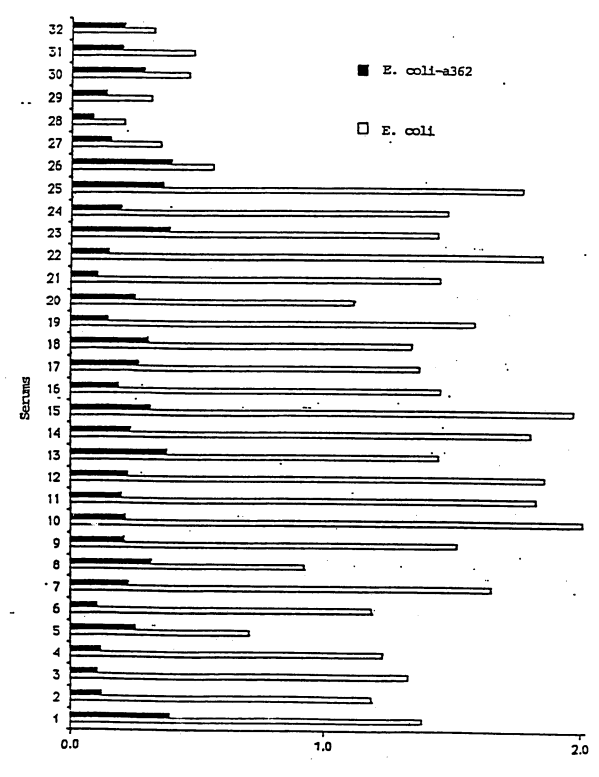
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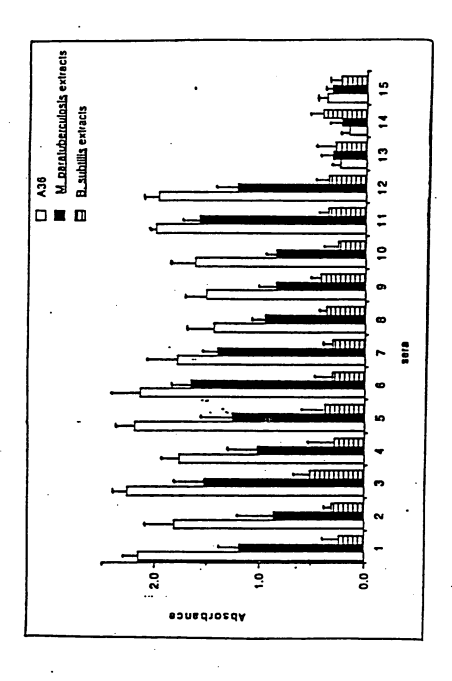
Figure 1 (1) 10 1 10 10 15 Figure 1 (2)

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Figure 2



Absorbance



F. 8. 3

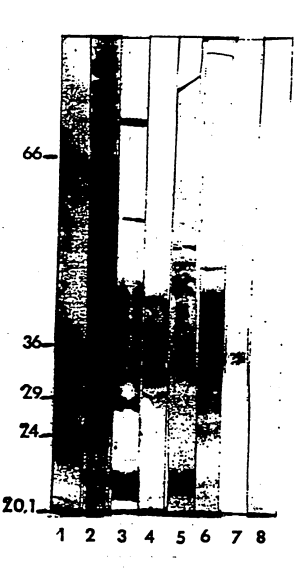


Figure 4

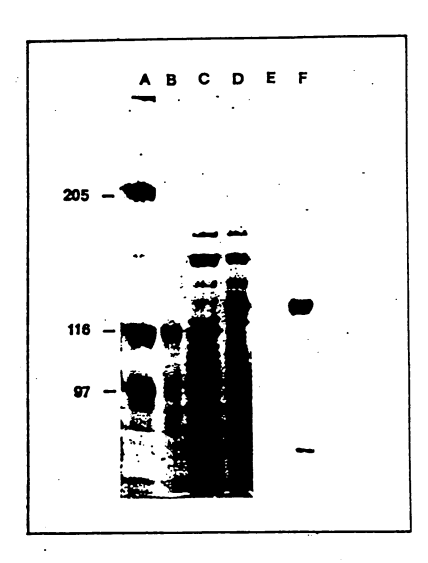


Figure 5



Figure 6

GAATTCCCGG		GTGGTCAGCA GCATTCGCCG CACCAGTCGT CGTAAGCGGC	CAGGXCTACG GTCCEGATGC	GGTCGCAGTA CCAGCGTCAT	CGGCGGTTAC GCCGCCAATG	09
GGCCAGGGCG	GCGCTCCGAC	CGGCGGTTTC GCCGCCAAAG	GGTGCCCAGC	CGTCGCCGCA	GTCCGGCCCG	120
CAACAGTCCG GTTGTCAGGC		CGCAGCAGCA GGGCCCGTCC GCGTCGTCGT CCCGGGCAGG	ACACCGCCCA TGTGGCGGGT	ACACCGCCCA CCGGCTTCCC CAGCTTCAGC TGTGGCGGGT GGCCGAAGGG GTCGAAGTCG	CAGCTTCAGC GTCGAAGTCG	180
CCGCYGCCCA		GGGATCGGAC CCCTAGCCTG	TCCGGTTCGG CGACCGCCAA AGGCCAAGCC GCTGGCGGTT	CGACCGCCAA GCTGGCGGTT	TTACTCCGAG AATGAGGCTC	240
CAGGCCGGTG	GECCAGCAGT	CCTACGGCCA	GGAGCCTTCT TCACCGTCTG CCTCGGAAGA AGTGGCAGAC	GGAGCCTTCT TCACCGTCTG CCTCGGAAGA AGTGGCAGAC	GGCCGACGCC CCGGCTGCGG	300
CGCCTAACGT		GCCTAGTCGG	GAACGTGCCC CAGAGTGACA	CAGAGTGACA GTCTCACTGT	CGGGTGGAGG	360
ACAACCGGGC		CGCCAGGCGC	GTGACCTCGT CACTGGAGCA	CAGGGTCGCG GTCCCAGCGC	TTCGCCCCGG AAGCGGGGCC	420
CGGTGGTGGC	CGGTGGTGGC ACTGGTCATC ATCGCCGCGG TCACGCTGAT CCAGTTGTTG GCCACCACCG TGACCAGTAG TAGCGGCGCC AGTGCGACTA GGTCAACAAC	ATCGCCGCGG TAGCGGCGCC	TCACGCTGAT AGTGCGACTA	CCAGITGITG GGTCAACAAC	ATCGCCAACA TAGCGGTTGT	480
GCGACATGAC	CGGCGCGTTG GGGAATTC GCCGCGCAAC CCCTTAAG	GGGAATTC CCCTTAAG				208
				•		
	.:	ħ	Fa. 7A			

Hg. 7A

300 360 420 480	GCCGACGCCC CGGCTGCGGG GGGTGGAGGA CCCACCTCCT TCGCCCCGGC AGCGGGGCCG	CACCGTCTGG GTGGCAGACC AGAGTCGCGT TCTCACTGTG TCCCAGCGCA CAGTTGTTGA GTCAACAACT	GAGCCITCIT CTCGGAAGAA AACGTGCCCC TTGCACGGGG TGACCTCGTC ACTGGAGCAG CACGCTGATC GTGCGACTAG	90 00 00 H& .	GGTGGTGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	TCCGGCCACC GCCTAACGTG CGGATTGCAC CAACCGGGCA GTTGGCCCGT GTTGGCCGGT CCACCACCGT CGACATGACC GGTGGTACCGGT
360		AGAGTGACAC TCTCACTGTG	AACGTGCCCC TTGCACGGG	CCTAGTCGGG GGATCAGCCC	CCCTGTCGCG	GCCTAACGTG CGGATTGCAC
300	CACCGTCTGG GCCGACGCCC GTGGCAGACC CGGCTGCGGG		GAGCCITCIT	CTACGGCCAG GATGCCGGTC	AGGCCGGTGG CCCAGCAGTC TCCGGCCACC GGGTCGTCAG	AGGCCGGTGG TCCGGCCACC
240	TACTCCGAGC ATGAGGCTCG	GACCGCCAAT	CCGGTTCGGC	CGTCGGCGGG GGATCGGACT CCGGTTCGGC GACCGCCAAT TACTCCGAGC GCAGCCGCCC CCTAGCCTGA GGCCAAGCCG CTGGCGGTTA ATGAGGCTCG		CGCGGCCCAA
180	GGCCCGTCCA CACCGCCCAC CGGCTTCCCC AGCTTCAGCC CCGGGCAGGG TCGAAGTCGG	CGGCTTCCCC	CACCGCCCAC GTGGCGGGTG	GGCCCGTCCA CCGGGCAGGT	AACAGTCCGC GCAGCAGCAG TTGTCAGGCG CGTCGTCGTC	AACAGTCCGC
120	TCCGGCCCGC	GTCGCCGCAG	GTGCCCAGCC	GGCGGTTTCG GTGCCCAGCC GTCGCCGCAG TCCGGCCCGC CCGCCAAAGC CACGGGTCGG CAGCGGCGTC AGGCCGGGCG	GCCAGGGCGG CGCTCCGACC CGGTCCCGCC GCGAGGCTGG	GCCAGGGCGG
9	GGCGGTTACG CCGCCAATGC	GTCGCAGTAC CAGCGTCATG	GCATTCGCCG CAGGCTACGG CGTAAGCGGC GTCCGATGCC		GTGGTCAGCA CACCAGTCGT	GAATTCCCGG CTTAAGGGCC

Fig. 7B

09	0	0		0	0	0	0	7
v	120	180	240	300	360	420	480	507
CGGCGGTTAC GCCGCCAATG	GTCCGGCCCG	GGGCCCGTCC ACACCGCCCA CCGGCTTCCC CAGCTTCAGC CCCGGGCAGG TGTGGCGGGT GGCCGAAGGG GTCGAAGTCG	TTACTCCGAG AATGAGGCTC	GCCGACGCCC CGGCTGCGGG	GGGTGGAGGA	TCGCCCCGGC AGCGGGGCCG	TCGCCGCGGT CACGCTGATC CAGTTGTTGA TCGCCAACAG AGCGGCGCCA GTGCGACTAG GTCAACAACT AGCGGTTGTC	·
GCATTCGCCG CAGGGCTACG GGTCGCAGTA CGTAAGCGGC GTCCCGATGC CCAGCGTCAT	CGGCGGTTTC GGTGCCCAGC CGTCGCCGCA GCCGCCAAAG CCACGGGTCG GCAGCGGCGT	CCGGCTTCCC	CGACCGCCAA GCTGGCGGTT	CACCGTCTGG GTGGCAGACC	CCTAGTCGGG AACGTGCCCC AGAGTGACAC GGATCAGCCC TTGCACGGGG TCTCACTGTG	GCCAGGCGCG TGACCTCGTC AGGGTCGCGT CGGTCCGCGC ACTGGAGCAG TCCCAGCGCA	CAGTTGTTGA GTCAACAACT	
CAGGGCTACG GTCCCGATGC	GGTGCCCAGC CCACGGGTCG	ACACCGCCCA TGTGGCGGGT	TCCGGTTCGG	CTACGGCCAG GAGCCTTCTT GATGCCGGTC CTCGGAAGAA	AACGTGCCCC TTGCACGGGG	TGACCTCGTC ACTGGAGCAG	CACGCTGATC GTGCGACTAG	·
GCATTCGCCG		CCCGGGCAGG	GGGATCGGAC	CTACGGCCAG GATGCCGGTC	CCTAGTCGGG	GCCAGGCGCG		GGAATTC
GTGGTCAGCA	GCGCTCCGAC CGCGAGGCTG	CAACAGTCCG CGCAGCAGCA GTTGTCAGGC GCGTCGTCGT	CCGCCGCCCA ACGTCGGCGG GGGATCGGAC TCCGGTTCGG CGACCGCCAA	CAGGCCGGTG GCCAGCAGTC CTACGGCCAG GAGCCTTCTT GTCCGGCCAC CGGTCGTCAG GATGCCGGTC CTCGGAAGAA	CCCTGTCGCG CCTAGTCGGG AACGTGCCCC AGAGTGACAC GGGACAGCGC GGATCAGCCC TTGCACGGGG TCTCACTGTG	CAACCGGGCA GCGGGCGCTC GCCAGGCGCG TGACCTCGTC AGGGTCGCGT GTTGGCCCGT CGCCGCGAG CGGTCCGCGC ACTGGAGCAG TCCCAGCGCA	GGTGGTGGCA CTGGTCATCA CCACCACCGT GACCAGTAGT	CGACATGACC GGCGCGTTGG GGAATTC GCTGTACTGG CCGCGCAACC CCTTAAG
GAATTCCCGG	GGCCAGGGCG CCGGTCCCGC	CAACAGTCCG	CCGCCGCCCA	CAGGCCGGTG	GCCTAACGTG	CAACCGGGCA GTTGGCCCGT	GGTGGTGGCA CCACCACCGT	CGACATGACC GCTGTACTGG

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GAA Glu	. TTC Phe	Pro	GGT	r GGT 7 Gly 5	CAC	G CAG	CAT His	TCG Ser	Pro 10	Glr	GGC Gly	TY	Gl?	TCG Ser 15	45
CAG Gln	TAC	Gly	GGT Gly	TAC Tyr 20	Gly	CAG Gln	GGC Gly	GGC	GCT Ala 25	Pro	ACC	GGC Gly	GGI Gly	TTC Phe 30	90
GCT	GCC Ala	CAG Gln	CCG	TCG Ser 35	CCG Pro	CAG Gln	TCC Ser	GGC Gly	CCG Pro 40	CAA Gln	CAG Gln	TCC Ser	GCG Ala	CAG Gln 45	135
CAG Gln	CAG Gln	GGC Gly	CCG Pro	TCC Ser 50	ACA Thr	CCG Pro	CCC Pro	ACC Thr	GGC Gly 55	Phe	CCC Pro	AGC Ser	TTC Phe	AGC Ser 60	180
CCG Pro	CCG Pro	CCC Pro	AAC Asn	GTC Val 65	GGC Gly	GGG	GGA Gly	TCG Ser	GAC Asp 70	TCC Ser	GGT Gly	TCG Ser	GCG Ala	ACC Thr 75	225
GCC Ala	AAT Asn	TAC Tyr	TCC Ser	GAG Jlu 80	CAG Gln	GCC Ala	GGT Gly	GGC Gly	CAG Gln 85	CAG Gln	TCC Ser	TAC Tyr	GGC Gly	CAG Gln 90	 270
GAG Glu	CCT Pro	TCT Ser	TCA Ser	CCG Pro 95	TCT Ser	GGG Gly	CCG Pro	Thr	CCC Pro	GCC Ala	TAA				306

Fig. 8

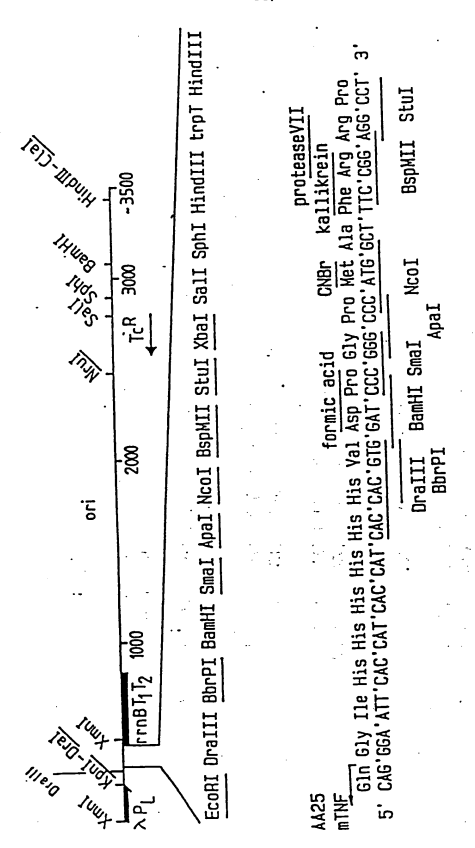


Figure 9a

Figure 9h

	45 - ' 'T'T'	K H	ပ္ခုပ္	ည်	i. Tr	H.K.	K.H.	ວິ ວ
	45 	NTA TAT	TAC	CTC	TTA AAT	TGT	TCA	GAG
	GCA	GTG	TGA	ACG	GGT	ອອວ	AAT TTA	ວອອ ອວວ
	39 	ນອນ ອນອ	ລວອ	GTG	GGA	CAA GTT	222 222	TTT AAA
	ນນນ	TCT AGA	TGG	AAG TTC	CCA	TGA	GCA	ອນນ
	33 T'GC A'C'G	CCA	CAC	ATG	GTA	GAG	GGA	CAT
	CAA GTT	TAA ATT	TAC	ACC	GGG GTA	TTC	GGA	ອອວ
	27 AAA TTT	AGA	AAA TTT	ACC TGG	GCA	AAA TTT	AGT	ວວອ ອອວ
	ACC	TAC	CAT	CTG	AGG	TCA	CCA	TCC
	21 	ACA TGT	TGA	GCA	AGA TCT	TAG	CCA	GGA
	TCA AGT	AAA TTT	TGT	GAC	TGA	AAG TTC	AAA	CGT
	15 CTC GAG	TAA	ວວອ	CAG	ອອອ	ATC TAG	AGC	CCA
Ĭ	GAT	ATA TAT	TGG	CAG	AAG	AAG	CGT	TCA
PMTNF HPH	6 6 6 7 7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	TTC	CTC	CAT	NTT TAA	GGT	CGT	CCA
pmT	TCC	AAA TTT	TAT ATA	GCA	AAA TTT	CAT	CCA	TCA
from:	3 AAT TTA	AAT TTA	AAT TTA	TGN	TTA	AAT TTA	AGC	CCA
<u>:</u>	Η,	46	91	136	181	226	271	316

ອອວ ວວອ	TTT AAA	AGA TCT	TAG	ACG	AGG TCC	ACT	TGA	AAC TTG
ອອວ	CTG	ATC	CAG	GAA	AGT	AAG TTC	TCC AGG	AGC
TAA	TGG	TAA	၁၁၅ ၁၁၁	AGT	GAG	CGA	CTC	CGA
AAG TTC	GCT.	GAT	TGG	AGA TCT	TGC	NGT TCA	ACG	TTG
AGT	CAA	ACA TGT	ອອວ ວວອ	CTC	CCA	CTC	TGA	ACG
TTA	ອອວ ວວອ	GAT	TTT AAA	GAA	TCC	AGG	ວວອ ອອວ	TGA
AGC	GAT	CCT	GAA	550	GTC	GAA	TGT	GAG CGG ATT TGA CTC GCC TAA, ACT
GCA	NTT NAA	CAG	ACA TGT	CAT	ວນນ	AAC TTG	GTT	ດ ດີດ ໂພນ 8
CAT	TTN	TTT	AAA TTT	ອອອ	TGT	TAA	GTT	GAG CTC
ວວອ	ATT TAA	GAT	GAT	TGA	TAG	AAA TTT	TCT	ວນອ
GAC	ອນນ	GAA	TCT	ACC	TGG	ATC	TTA	ອນອ
GT'C CAG	ອນນ	AGA TCT	၁၁၅ ၁၅၁	၅၅၅ ၁၁၁	CGA	ອນນ	GTT	ATC
AGA TCT	GCT	ATG	AAG TTC	GGT	ອນອ	CCA	TTC	CAA GTT
TCT AGA	TCC	၁၁၅ ၁၅၁	CAG	GGT	TAG	CTG	CCT	GGA
550 505	AGT TCA	TGG	ACG	ອນອ	ນນນ	GAA	000 000	GTA
361	406	451	496	541	586	631	919	721

ນນ	CGT	ATG	TAA ATT	AAT TTA	AGA TCT	AAT T'FA
CCA	TTG	AAT TTA	TAN	CAA GTT	CGT	CGT
CTG	TTT	CAT TCA GTA AGT	CAA GTT	GAC	<u>ອ</u> ອອ	ນອນ ອນອ
aaa T't	ອອວ ວວອ	CAT	CTT	TCT CAT	AGA TCT	TCT AGA
CAT	NTG	TY.	TG	TCT	GTC	TTT AAA
ອນອ	ວວອ ອອວ	TAA	TA	TTT TGA TAA AAA ACT ATT	CCA CTG AGC GGT GAC TCG	TTT AAA ''t)
၅၅၁ ၁၁၅	TGA ACT	TT(AA(TGA	TGA	CTG	AGA TCC TTT TCT AGG AAA Figure 9b (con't)
GAC	TCC	TTT	ລວວ	TTT AAA	CCA	AGA TCT Figure
GGG CAG CCC GTC	CCA	TTA	TAA	CCT	GTT CAA	TTG AAC
ນນນ	AGG	TGT	CAA GIT	GAT	TTC	TTC
ອນນ	ngn TCT	TTT AAA	AGA TCT	GAA	GTT	ATC TAG
CCA	AGC	CTC	ATG	GGT	ACG TGA TGC TGC TGC	AGG ATC TCC TAG
GAG	T'TA AAT	AAA T'T	CTC	CTA	ACG	CAA GTT
ນນນ	AAA TTT	TAC	ວວວ	GAT	TTA AAT	GAT
ອນນ ນອອ	atc Tag	T'TC	TAT ATA	AAG TTC	ວວວ	AAA TTT
166	811	856	901	946	991	1036

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TTG	TGG	ອອວ	ATA TAT	CGA	GGA	ອອວ ວວອ
GGT	AAC TTG	GTA	TAC	TGG	ACC TGG	ACA TGT
GGT	GGT CCA	AGT	GCC TAC	CAG	GTT	CAC
AGC GGT OTCE OCCE	GAA	TCT	ACC TGG	TGC	ATA TAT	GTG
ACC TGG	TCC	CCT	AGC TCG	TGC	ACG TGC	TTC AAG
GCT	TTT AAA	TGT	TGT	ອນວ	AAG TTC	(2, uo2) 222 222 222 232 203 203 203 203 203 203
ACC TGG	TCT AGA	TAC	CTC	AGT TCA	CTC	(3, wo) CCC BBB
ACC ACC	AAC	AAA TTT	GAA	ACC TGG	GGA CCT.	CTG AAC GAC TTG Figure 9b
AAA TTT	ACC	ACC TGG	CAA	GTT	GTT	
AAA TTT	GCT	GAT	CTT	CCT	ນນນ	ນ ນ ນ
AAC TTG	AGA	GCA	CCA	aat Tta	TAC	GTC
GCA	TCA	AGC	CCA	GCT	TCT	ນ ອນອ
CTT	GGA	CAG	AGG	TCT AGA	GTG	GCA
CTG	ນນອ	CAG	GTT CAA	ອນອ	GTC	ອນນ ນອອ
CTG		CTT	GTA	CCT	TAA	TAA
1081	1126 TTT AAA	1171	1216	1261	1306	1351 TAA ATT

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ນນນ	GGA	GAG	ວນນ	GTC
ACA TGT	ອນນ	CAC G'IG	TGT ACA	CTC
CCT	AAA TTT	ນອນ ອນອ	TCC AGG	ATG TAC
ATA TAT	GAG	AGA TCT	TAG ATC	GTG
AAC GAC CTA CAC CGA ACT GAG ATA TTG CTG GAT GTG GCT TGA CTC TAT	CGA AGG (GCT TCC (CGG AAC AGG	CTG GTA TCT TTA GAC CAT AGA AAT	TTT GTG ATG AAA CAC TAC
ACT	CGA	AAC TTG	TCT	ATT TAA
CGA	TCC	ວວຣ ອອວ	GTA	M GCG TCG I CGC AGC Figure 9b (con't.
CAC	GCT	GGT	CTG	GCG CGC
CTA	CAC	CAG	ອນອ	TGA ACT
GAC	CGC	ນນນ	AAA CGC (TILL CCC C	ACT
		AAG	ວວວ ອອອ	CTG
GGA GCG CCT CGC	AGA TCT	GGT	AGG	CCT
GGA	TTG	TCC	TCC	CCA
CTT GAA	GCA	GTA	GCT	TCG
CAG	TGA	CAG	GGN	GTT
1396 CAG CTT GTC GAA	1441 TGA GCA ACT CGT	1486	1531 GGA GCT TCC CCT CGA AGG	1576
				•

1621	AGG	ນນນ	ນນິນ	GAG	CCT	ATG	GNA	AAA TTT	ນນຸນ	CAG	CAA	၅၃၅ ၁၅၁	ອນນ ນອອ	CTT	TTT
. 1666	ACG	GTT	CCT	ນນນ	CTT	TTG	CTG	ອອວ	TTT TGC NAN ACG	TGC	TCA	CAT	GTT	CTT	TCC
1711	TGC	GTT	ATC	CCC TGA 1 GGG ACT 1	TGA	rTC \AG	TGT	TGT GGA ACA CCT	TAA ATT	ນນຸນ ອນນ	TAT ATA	TAC	ອນອ ນອນ	CTT	TGA
1756	GTG	AGC	TGA	TAC CGC ATG GCG	၅၁၅ ၁၅၁	rcg Agc	້ວຍຍ	CCG CAG	ນອອ ອນນ	AAC TTG	GAC	CGA	ວອວ ອວອ	CAG	CGA
1801	GTC	AGT	GAG	CGA	GGA	AGC	GGA	GGA AGA CCT TCT	ນອນ ອນອ	CTG	ACT TGA	TCC	ນ ອນອ	TTT	CCA
1846	GAC	TTT	ACG	AAA TTT	CAC		AAC TTG	CGA	AGA TCT	CCA GGT	TTC		TTG	TTG	CTC
1891	AGG	TCG	CAG	ACG TGC	TTT	TGC	AGC TCG	AGC AGC AGT TCG TCG TCA	AGT TCA	ອນອ	TTC		TTC	GCT	ອນອ ນອນ

AGC	GTG	TGG	ນນນ	GAG	CGA	GAC
ອອນ ນນອ	ອອອ ວວວ	TGC	TTT	TTG	GGT	GCA
ອອອອ	GCA	ງງງ	TGG	TTC	TCA	GAG
AAC TTG	TGC	TGC	GGT TGG	CAA	CAT	999
ອນນ	TCA	ນຄນ ອນອ	AAG TTC	CTC GAG	TTC	ညည
TAA	CGA	ອອວ ວວອ	TGT TCT GCC AAG ACA AGA CGG TTC	TGG	ອນນ ນອອ	ACC GCG ACG CAA CGC GGG GAG GCA GAC TGG CGC TGC GTT GCG CCC CTC CGT CTC
CAG	GCA	TGC	TCT	GAT	999 9999	ACG TGC
AAC	GGA	AGA TCT	TGT	ATT	ອອວ	3 3 3 3 3 3 3 3
GCT	ACA	ນນນ	ATA TAT	GCA AGA CGT TCT	GGT	ACC TGG
CAT TCT GTA AGA	acg	TGC	TGG	GCA	CGA	TGC
CAT	TCC TCA AGG AGT	CAA CGC GTT GCG	CGA	TTC TCC AAG AGG	TAG	GCT CCA CGA GGT
ATT	TCC	CAA	ACG	TTC	CGT GCA	GCT
GTG	ລວວ	ACC TGG	ລລອ	CAG	atc Tag	ນນນ
TCG	ອອວ	AGG	TGG	TCA	TGA	ອນນ
GTA	CTA GAT	ອອວ	AGA TCT	CAT GTA	TGG	GGT
1936	1981	2026	2.071	2116	2161	2206

	CGC CGT GAC GAT CAG CGG TCC AGT GCG GCA CTG CTA GTC GCC AGG TCA	AGC CGC GAG CGA TCC TTG AAG CTG TCG GCG CTC GCT AGG AAC TTC GAC	CTG CCT GGA CAG CAT GGC CTG CAA GAC GGA CCT GTC GTA CCG GAC GTT	GGA AGC GAG AAG AAT CAT AAT GGG
ATG TTA	AAT TTA	AAG	TAC	000
990	ATA TAT	GGT	ATC TAG	99 2
) () ()	ອວວ ອວວ	GCT	GTC GTC CAG CAG	GAT
0 0 0 0 0	ອນນ	TAG	GTC	<u>ອ</u> ອອອ
ATC	CGA	AGT	ATG	CAT
	ອນ ນອນ	CGA	CTG	ອຍອ ວິດ
TTC	GCT	GAT	TCC	555 555 555
1622	2296	2341	2386	2431

CGC GTG GGC CAT GGC GTT GGG GTG GTT GGG GTG GTT GTG GTT GGG GTG GTT GTG GTT GTG GTT GGG GTG GTT GTG GTT GTG GTT GGG GTG GTT GTG GTT GGG GTG GT	ອອວ ວລອ	ນ ນ ນ ນ	ອນນ ນອອ	ອນອ ນອນ	TGC	AAG	GAC
GAA GGC CAT CCA GCC TCG CGT CGC GAA CGC CAG CTT CCG GTC GTC GTT CCG GTA GGG CTT GCG GTC GTC GTG CTT GCG GTC GTC	GTA			CG'T GCA	ອນອ ນອນ		GCT
GAA GGC CAT CCA GCC TCG CGT CGC GAA CGC CAG CTT CCG GTC GTC GTT CCG GTA GGG CTT GCG GTC GTC GTG CTT GCG GTC GTC		CTT	AGC TCG	CAT GTA		AGT TCA	GGA
GAA GGC CAT CCA GCC TCG CGT CGC GAA CGC CAG CTT CCG GTA GGT CGG AGC GCA GCG CTT GCG GTC CAG CGC GTC GGC CGC CAT GCC GGC GTA TTA CCG GTC GCG CAG CCG GTA CGG CTA TTA CCG GAA ACG TTT GGT GGC GGG ACC AGT GAC GAA GGC CTT TGC AAA CCA CCG CCC TGG TCA CTG CTT CCG GTG CAA GAT TCC GAA TAC CGC TTC GCT GTC CGG GTG CAA GGC CTT ATG GCG TTC GCT GTC CGG CCA GTT CTA AGG CTT ATG GCG TTC GCT GTC CGG CCG CAC CTG TTT CGC GAG TTC GCT GTC CTG CGG CAC CTG TTT CGC CAG GAG CGG CTT TTA CTG CGG CAC CTG TCC TAC GAG TTG CAT TTA CTG TGC GTG GAC GAC GAG CTC TTT TTA CTT TGC GGC GAC GAC GAT GTT CTT TTA CTT TGC GGC GAC GAC CAT GAT AAA GAA GCC GTG GAC GAC CAT CTA TTT CTT TGC GGC GAC CAT TTT CTT TGC GGC CAC CTG CAT CTT TTA CTT TGC GGC GAC GAC CAT CAT CTT TTA CTT TGC GGC GAC GAC CAT CAT CTT TTA CTT TGC GGC CAC CGC CCC CCC CCC CCC CCG		CTG	TTG	GAT		GAC	GAA
GRA GGC CAT CCA GCC TCG CGT CGC GAA CGC CTT CGC GTT CGC GTT CCG GTT CGG AGC GCA GCG CTT GCG GTT GCG GTA CGG CTT TTA GGT GGG GTG GTA CGG CTA TTA GGT GGG GTG CGG CTA TTA GGT GGC GGG ACC AGT GAC GTA CTT TGC AAA CCA CCG CCC TGG TCA CTG CTT GCT GTT CAC GTT CTA AGG CTT ATG GCG TTC GCT GTC GTC GTT CGA GTT CGC GTT TTA CGC CAG GGG CTT TTA CGC CAG GAG CGG CTT TTA CGC CAG GAG CGG CTT TTA CGG CAG GAG CGG CCC CCA CGC CCC CCA CGG CAG CGC CCA CCA		ອນນ ນອອ	ອນນ	ອອວ ວວອ	GAC	GAA CTT	ລອອ
GAA GGC CAT CCA GCC TCG CGT CGC GAA CTT CCG GTA GGT CGG AGC GCA GCG CTT CAG CGC GTC GGC CGC GTA CGG CTA GTC GCG CAG GCG GTA CGG CTA GAA ACG TTT GGT GGC GGG ACC AGT GAC CTT TGC AAA CCA CCG CCC TGG TCA CTG GTG CAA GAT TCC GAA TAC CGC AGG CGT CAC GTT CTA AGG CTT ATG GCG TTC GCT GCT CCA GCG AAA GCG GTC CTC GCC GAA CGA GGT CGC TTT CGC GAG CGG CTT CGG CAC CTG TCC TAC GAG CGG CTT CGG CAC GTG TCC TAC GAG GAG CGG CGG CAC CTG TCC TAC GAG GTG CTT CGG CAC GTG TCC TAC GAG TTG CAT GAT TGC GGC GAC GAG ATG CTC AAC GTA TGC GGC GAC CAC CAC CCC CCC CGC		aat T'ea	GAA	CAG	aat Tta	ann Ttt	CCA
GRA GGC CAT CCA GCC TCG CGT CGC CTT CCG GTA GGT CGG AGC GCA GCG GAG CGC GTC GGC GGC GTA CGG CCG GTC GCG CAG GCG GTA CGG CCG GTC GCG CAG GCG GTA CGG CCG GTA ACG TTT GGT GGC GCG CCC TGG TCA GTG CAA ACG TTT GGT GGC GCG CCC CTT TGC AAA CCA CCG CCC TGG TCA GTG CAA GAT TCC GAA TAC CGC AAG GCT CCA GCG AAA GCG GTC CTC GCC CGA GGT CGC TTT CGC CAG GAG CGG CGG CAC CTG TCC TAC GAG TTG CAT TGC GGC GAC GAT AGT CAT GCC CCG	GAA	GAT		CGA	GAA CTT	GAT	
GAA GGC CAT CCA GCC TCG CTT CCG GTA GGT CGG AGC CAG CGC GTC GGC GCG GTP GAA ACG TTT GGT GGC GGG CTT TGC AAA CCA CCG CCC GTG CAC GTT TCC GAA TAC GTG CAA GGT CTT ATG GCT CCA GCG AAA GCG GTC CGA GGT CGC TTT CGC CAG GCT CCA GCG AAA GCG GTC CGA GGT CGC TTT CGC CAG TGC GTC TTT CGC CAG TGC GTC GAC AGG ATG CTC TGC GGC GAC GAT AGT CAT	ອນອ	ອນນ ນອອ	AGT TCA	AAG TTC		CAT	ວນອີ
GAA GGC CAT CCA GCC TCG CTT CCG GTA GGT CGG AGC CAG CGC GTC GGC GCG GTP GAA ACG TTT GGT GGC GGG CTT TGC AAA CCA CCG CCC GTG CAC GTT TCC GAA TAC GTG CAA GGT CTT ATG GCT CCA GCG AAA GCG GTC CGA GGT CGC TTT CGC CAG GCT CCA GCG AAA GCG GTC CGA GGT CGC TTT CGC CAG TGC GTC TTT CGC CAG TGC GTC GAC AGG ATG CTC TGC GGC GAC GAT AGT CAT	CGT		ACC	ອນອ ນອນ		TTG	300 200 100 100 100 100
GAA GGC CAT CCA GCC CTT CCG GTA GGT CGG GTC GCG CGG GTT GTT GGT GGC CTT TGC AAA CCA CCG GTG CAA GCG CTT CTA AGG CTT GCA GCG TTT CGC CGA GGT CGC TTT CGC CGA GGT CGC TTT CGC CGG CAC CTG TCC TAC GCG CAC GTG GAC ATG TGC GGC GAC GAT AGT	TCG		ນນນ	TAC		GAG	CAT GTA
GAA GGC CAT CTT CCG GTA CAG CGC GTC GTC GCG CAG GAA ACG TTT CTT TGC AAA GTG CAA GAT CAC GTT CTA GCT CCA GCG CGA GGT CGC CGA GGT CGC CGG CAC CTG GCC GTC GAC TGC GGC GAC				GAA	້ນຄວ		AGT
CTT CCG GTP CTT CCG GTP CAG CGC GTC GTC GCG CAG GAA ACG TTT CTT TGC AAA GTG CAA GAT CAC GTT CTA GCT CCA GCG CGA GGT CGC CGA GGT CGC CGG CAC GTG GCC GTG GAC	CCA	ອວວ	GGT	TCC	aaa Ttt	TCC	GAT CTA
GAA GGC CTT CCG GTC GCG GTC GCG CTT TGC GTG CAA CGC GTT GCT CCA CGC GTC		GTC	TTT AAA	GAT	ວອວ		GAC
GRAA CTT GRA GTG CAC GCT GCT CGA CGA CGA		ອນອ	ACG TGC	CAA GTT			ອນນ ນອຍ
• _		CAG	GAA CTT	GTG	GCT	ວວອ ອອວ	
	2476	2521	2566				

			21	/27		
NTG	GTT	ອອວ ວວອ	GAA	ATC	ອອວ ວວອ	GAC
CTT	ອອວ	၁၁၁	ອອວ	ອອອ	ອນນ ນອອ	CAG
TCC	GAG	GAT	CAC	T'TC AAG	TGT	CCA
CTC	GTT	GGA	ACC	ATC	ACC TGG	GAT
ACG	TAG	CAA	CAT	CCG ATC GGC TAG	ອນອ ນອນ	GAG
TCG	TAG	ATG TAC	CAC	AGC TCG	AAC TTG	GTA
ຍຍວ	CAG	TGC	TGC	ລອວ	AGC	000 000 000
CAT	ອອວ ວວອ	TGG	ອອລ	GTG	ອອວ	GCG TCC GGC CGC AGG CCG Figure 9b (con't)
ນນນ	GCA	GNA	ວນວ	GNA	ອນນ	GCG CCC Figure
CAA GTT	GAA	AAG	CAC	ອອອ	ATA TAT	GAT
TCT AGA	TAG	ອນອ	ອນນ	GAG	GAT	CAC
ນນນ	CAT	ອນອ	ອອອ ນນນ	CAT	ອນນ	ອນນ
GNA CTT	CTG	ອນອ	TCC AGG	GCT	GTC	ອອວ
GTT CAA	CTC	CAC	CAG	AGC	GAT	GAT
TGG	CGA	GAG	CAA	ACA TGT	GGT	GGT
2791	2836	2881	2926	2971	3016 GGT CCA	3061

_	_		_	_
7	7	/	7	7

TAG	TGC	ອນອ ນອນ	GAT	ອອນ ວວອ
AAG	CAG	rag atc	GAC	TAT ATA
TCC AAG AGG TTC	GGA	ATA TAT	ATG TAC	ဗိဗ္ဗာ ၁၁၅
ອນນ ນອອ	GTC	CGC ATA 1 GCG TAT 2	GGA	CAA GTT
AGT	300	CAA GTT	GTC	AAC TTG
GAT	GCC AAA (CCC TIT	CAT	GAT GCT CTA CGA	CAT
GTC	ອອວ ວວອ	TTG	GAT	GGG CAG TAC CGG (CCC) GCC (Figure 9b (con't)
GTA	၂၅၁ ၁၁၅	AAA TTT	ອນນ ນອອ	TAC ATG
ອນອ	ວຼວວ	TAG	ACT	CAG GTC Figure
GAT	TGG	GCA	GTG ACT	ວວອ ອອວ
CAT	GAC	TGC C	ATA TAT	ည်း
ອນອ	CAG GAC GTC CTG	ນນນ	GCC ATA (CGG TAT	GAG
GGT	GAG CTC	AAC TTG	CAC	CAA
TĠT ACA	AGC TCG	GAG	CAG	ວອອ ອນວ
ນນນ	CGA	TCC GAG	TAG	ATC TAG
3106 GGG 7 CCC 7	3151	3196	3241	3286

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933

845

Total number of bases is: DNA sequence composition:

name: NPMTNFMPH.

2 OTHER; Sequence

NTT TAN	CTG	AAC TTG	
GAT GAG CGC ATT CTA CTC GCG TAA	TAA	TCA AGT	
GAG	ACT GCG TTA GCA ATT TGA CGC AAT CGT TAA	GAT GAT AAG CTG CTA CTA TTC GAC	
GNT	GCA	AAG TTC	
GAC	TTA AAT	GAT	
GAT GAC CTA CTG C	ວອວ	GAT	••
GAG	ACT	ATC TAG	
<u>ອ</u> ອວ ວວອ	CTG	CTT	
GGT	TGC	AAG	
CAG GGT GAC GGT GCC GAG GTC CCA CTG CCA CGG CTC	CAT ACA CGG GTA TGT GCC	TTA	ï
GGT	ACA TGT	ACC GCA TGG CGT	
CAG	CAT	ACC TGG	
ATC TAG	TTT AAA	ACT TGA	ATT
AGC ATC TCG TAG	AGA TC'I	TAA	AGA ATT TCT TAA
TAC	GTT	TGA	ATG
3331	3376	3421	3466

Met Val Arg Ser Ser Ser Gln Asn Ser Ser Asp Lys Pro Val Ala 1 His Val Val Ala Asn His Gln Val Glu Glu Gln Gly Ile His His 16 31 His His His Val Asp Pro Gly Pro Met Ala Phe Arg Arg Pro Leu Glu Phe Pro Gly Gly Gln Gln His Ser Pro Gln Gly Tyr Gly 46 Ser Gln Tyr Gly Gly Tyr Gly Gln Gly Gly Ala Pro Thr Gly Gly 61 Phe Gly Ala Gln Pro Ser Pro Gln Ser Gly Pro Gln Gln Ser Ala 76 Gln Gln Gln Gly Pro Ser Thr Pro Pro Thr Gly Phe Pro Ser Phe 91 Ser Pro Pro Pro Asn Val Gly Gly Ser Asp Ser Gly Ser Ala 106 Thr Ala Asn Tyr Ser Glu Gln Ala Gly Gly Gln Gln Ser Tyr Gly 121 136 Gln Glu Pro Ser Ser Pro Ser Gly Pro Thr Pro Ala

GGG	ccc	GAA	CIT	GAC	GAA	CTC	GCC	GTC	GTA	GCT	GGC	TTC	CTC	GTC	45
GGT	CCA	CAG	CGC	CCG	CAT	CGC	TTC	CAG	GTA	TTC	GCG	CAG	CAT	GGT	90
GCG	GĊG	CCG	GCC	CGC	CGG	CAC	GCC	GTG	GTC	GGC	GAG	TTC	GTC	GGT	135
GTT	CCA	GCC	GAA	ccc	GAC	GCC	GAG	GCT	GAC	CCG	GCC	GCC	GGA	CAG	180
ATG	GTC	AAG	GGT	GGC	AAT	ACT	TTT	CGC	CAG	CGT	GAT	CGG	GTC	GTG	225
TTC	GAC	CGG	CAG	GGC	CAC	CGC	GGT	GGA	CAG	CCG	CAC	CCG	CGA	GGT	270
GAC	GGC	ACA	GGC	CGC	GCC	CAG	ACT	GAC	CCA	CGG	GTC	CAG	GGT	GCG	305
CAT	GTA	GCG	GTC	GTC	GGG	CAG	CGA	CGC	GTC	GCC	GGT	GGT	CGG	GTG	360
CGC	GGC	CTC	CCG	CIT	GAT	CGG	GAT	ATG	CGT	GTG	TTC	CGG	CAC	GTA	405
GAA	GGT	CGC	AAA	CCC	GTG	GTC	GTC	GGC	AAG	CTT	CGC	GGC	CGC	AGC	450
CGG	AGA	GAT	GCC	ACG	GTC	GCT	GGT	GAA	AAG	CAC	AAG	CCC	GTA	ATC	495
CAT	GCA	GTG	AAT	TAG	AAC	GTG	TTC	TAC	CTC	TGC	GGG	GCA	AGC	TGT	540
CGT	GAT	ACG	GAC	CGT	CTC	GCC	GCG	CGG	TCG	TCT	GCG	AAG	ccc	GCG	585
GGC	AAG	CCA	ATG	GCG	ACG	GCA	CCG	GCC	GTC	GCA	CGT	GCG	CTA	GCG	630
TGG	GTG	ATC	GAC	CGT	GTC	GCT	CGC	GCA	GTG	ACG	CGC	CTG	CAA	GCA	67 <u>,</u> 5
CCG	CGT	CGC	ATC	GCA	ACC	GTG	GCG	ccc	GCT	CGG	CAC	TAA	AAG	GCA	720
GTG	GAA	GCA	ACA	GGA	GGA	GCC						Gly			765
												GGC			810
							Asp					CTC Leu			855
												GCC Ala			900
												CTC Leu			945

Figure 11

GGT ATC GGC GGC GGC GGT GAC GCC GGC ACC GCC GTC GTG GTG G1 Ile Gly Gly Arg Ala Gly Asp Ala Gly Thr Ala Val Val Val 70 75 80

GCG CTG CTG GCC GCG CTG CTC GCC GGG CTG GGC CTG CCC AAG 1035
Ala Leu Leu Ala Ala Leu Leu Ala Gly Leu Gly Leu Leu Pro Lys
90
95

GCC AAG AGT TAT GTG GGC GTG GTC GCG GTC GCG GTG CTC GCC 1080 Ala Lys Ser Tyr Val Gly Val Val Ala Val Ala Val Leu Ala 100 105 110

GCG CTG CTG GCC ATC ACC GAG ACG ATC AAC CTG CCC GCC GGT TTC 1125 Ala Leu Leu Ala Ile Thr Glu Thr Ile Asn Leu Pro Ala Gly Phe 115

GCG ATC GGC TGG GCG ATG TGG CCG CTG GTG GCG TGC GTG GTG CTG 1170
Ala Ile Gly Trp Ala MET Trp Pro Leu Val Ala Cys Val Val Leu
130 135 140

CAG GCG ATC GCC GCG GTG GTC GTG GTC CTG CTG GAC GCC GGG GTG 1215 Gln Ala Ile Ala Ala Val Val Val Leu Leu Asp Ala Gly Val 145 150 155

ATC ACG GCG CCG GCG CCG CCC AAG TAC GAC CCC TAC GCG CAG 1260

Ile Thr Ala Pro Ala Pro Arg Pro Lys Tyr Asp Pro Tyr Ala Gln

160 165 170

TAC GGC CAA TAC GGG CAA TAC GGC CAG TAC GGG CAA CAG CCC TAC 1305
Tyr Gly Gln Tyr Gly Gln Tyr Gly Gln Gln Pro Tyr
175

TAC GGT CAG CCG GGC GGT CAG CCC GGG GGC CAG CCG GGT GGT CAG 1350
Tyr Gly Gln Pro Gly Gly Gln Pro Gly Gly Gln Pro Gly Gly Gln
190
195
200

CAG CAT TCG CCG CAG GGC TAC GGG TCG CAG TAC GGC GGT TAC GGC 1395
Gln His Ser Pro Gln Gly Tyr Gly Ser Gln Tyr Gly Gly Tyr Gly
205
210
215

CAG GGC GGC GCT CCG ACC GGC GGT TTC GGT GCC CAG CCG TCG CCG 1140
Gln Gly Gly Ala Pro Thr Gly Gly Phe Gly Ala Gln Pro Ser Pro
220 225 230

CAG TCC GGC CCG CAA CAG TCC GCG CAG CAG CAG GGC CCG TCC ACA 1485
Gln Ser Gly Pro Gln Gln Ser Ala Gln Gln Gln Gly Pro Ser Thr
235 240 245

Figure 11 (con't 1)

1839

27*2*7

										CCG Pro					1530
										AAT Asn					1575
										CCT Pro					1620
				GCC Ala		CGT	GCC	CTG	TCG	CGC	CTA	GTC	GGG	AAC	1665
GTG	ccc	CAG	AGT	GAC	ACG ·	GGT	GGA	GGA	CAA	CCG	GGC	A GC	GGG	CGC	1710
TCG	CCA	GGC	GCG	TGA	CCT	CGT	CAG	GGT	CGC	GTT	CGC	ccc	GGC	GGT	1755
GGT	GGC	ACT	GGT	CAT	CAT	CGC	CGC	GGT	CAC	GCT	GAT	CCA	GTT	GTT	1800

Figure 11 (con't 2)

GAT CGC CAA CAG CGA CAT GAC CGG CGC GTT GGG GAA TTC

L CLASSIFICATION OF SUBJE	CT MATTER (If several classification sys	mhois apply, indicate ail) ⁶		
According to International Patent	Classification (IPC) or to both National Cla	estification and IPC		
Int.C1. 5 C12N15/33 C07K13/00	1; G01N33/569;	A61K39/04; C12N1/21;		LK39/395 2N5/10
IL FIDLDS SEARCHED				
# F/2233 30-31-25	Misimum Documen	station Searched		
Chestification System		Jestification Symbols		
CALLERON DIA				
Int.Cl. 5	CO7K; C12P;	C12Q ;	A61K	
	Decementation Searched other to to the Extent that such Documents a	han Mainam Documentation re Incinied in the Fields Sear	del ⁸	
III. DOCUMENTS CONSIDERE	ED TO BE BELEVANT ⁹			
Category * Circuites of De	sceneri, 11 with Indication, where approprie	es, of the relevant passages 12		Edwar to Calm No.13
A WO,A,8 FOUNDAT	903 892 (WISCONSIN ALUM ION) 5 May 1989 e 4, paragraph 2 e 6, paragraph 4 - page	NI RESEARCH		5
MCFADDE see abs	808 456 (J. HERMAN-TAYL N) 3 November 1988 tract e 27, paragraph 2	OR & JJ.		1,5, 17-24
		-/	•	
	·			
"I" carrier decement but pair filing date "I" focument which may then which is cled to establish citation or other special a clear other special a charactering to an other special.	meral size of the art which is not calar relevance lished on or after the interestional over doubts on priority cisim(s) or in the publication date of smother reason (as specified) a cral discipulary, see, exhibition or r to the interestional filling date but	"I" later document publish or priority data and not cited to understand the invention. "I" document of particular cannot be considered as inventive at inventive at inventive at the cannot be considered a document of considered passes, such combined passes, such combined in the art. "A" document member of the	the entitlet with the principle or theory relevance; the cixis and or cannot be a property of the cixis and the cixis are an invest with one or more a page being obvious to	ne gypection we y underlying the med invention med invention tre step when the ther such deca- t a person shilled
IV. CEXTEFICATION				
Date of the Actual Completion of	the International Search MAY 1992	Date of Mailing of this 2 3, 0		ch Report
International Searching Authority EUROPE	ZAN PATENT OFFICE	Signature of Authorized THIELE U.	Officer	~.~.

Parts PCT/ISA/210 (accord about) (Jamesy 1963)

	· Interestional Application No	
m nocime	NTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	
	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim Na.
Category *		
A	JOURNAL OF CLINICAL MICROBIOLOGY vol. 25, May 1987, WASHINGTON D.C., US pages 796 - 801; J. J. MCFADDEN: 'Crohn's Disease-Isolated Mycobacteria Are Identical to Mycobacterium paratuberculosis, as Determined by DNA Probes That Distinguish between Mycobacterial Species' see abstract see page 798, left column, paragraph 5 - right column, paragraph 1 see page 799, right column, line 37 - page 800, line 5	5
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03-11-88	EP-A- EP-A-	0288306 0356450	26-10-88 07-03-90
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